Page 1 of 1

STIC-EIC1600/2900

From: SEAN BASQUILL [sean.basquill@uspto.gov]
Sent: Friday, March 06, 2009 10:48 AM

To: STIC-EIC1800/2900

Bubject: Search Request, Case/Application No.: 10560012

Requester: SEAN BASQUILL (P/1612) Art Unit: GROUP ART UNIT 1612

Case/Application number: 18560012
Priority Filing Date: 10 June 2003
Format for Search Results: Email
Meaning of unusual acronyms or initialisms:

Identify the novelty:

Chains 29, as attached, recites approximately four-and-a-half pages of discrete chemical compound names of species within a genus ac claimed. I would like to know if any of the compounds were known in the art prior to June 10, 2005.

Additional comments:

ONLY the chemical compounds as recited in Claim 29 (which yes, goes on for five pages) are of concern here.

Attachment: Yes (10560012 Claim 29.mlf)

362000

=> file registry

FILE 'REGISTRY' ENTERED AT 11:02:00 ON 12 MAR 2009
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Property values tagged with IC are from the ZIC/VINITI data file

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STRUCTURE FILE UPDATES: 11 MAR 2009 HIGHEST RN 1119363-64-2 DICTIONARY FILE UPDATES: 11 MAR 2009 HIGHEST RN 1119363-64-2

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TSCA INFORMATION NOW CURRENT THROUGH January 9, 2009.

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

Uploading L26.str

chain nodes:
8 9 10 11 15 20 29 30 31 32 33 35 37 38 39 41 49 50 51 52 53

54 55 56 57 58 66 67 68 69 70 71 72 76 77 78 79 87 88 89 90 91 92 93 94 95

99 100 101 102 103 104 106 107 108 109 110 111 112 113 115 116 117 118 120 121

122 123 124 125 135 136 137 138 139 140 143 144 145 146 150 153 154 156

ring nodes :

Connectivity:

```
1 2 3 4 5 6 22 23 24 25 26 27 43 44 45 46 47 48 60 61 62 63
64 65 73 74 75 81 82 83 84 85 86 126 127 128
chain bonds :
1-20 2-32 3-31 4-8 5-35 6-33 9-153 9-154 15-156 22-30 23-39 24-37 25-29
26-38 27-41 43-54 44-50 45-49 46-52 47-53 48-51 54-55 54-56 55-58 56-57
60-70 61-76
62-69 63-66 64-78 65-77 66-67 66-68 70-71 70-72 72-73 79-84 81-88 82-94
83-95 85-87
86-93 88-90 88-89 89-92 90-91 100-101 101-103 102-103 102-104 106-108
106-107 109-110
110-111 110-113 111-112 115-150 116-117 117-118 117-120 121-122 121-123
124-125 135-136
136-137 138-139 139-140 143-144 143-145 143-146
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6 22-23 22-27 23-24 24-25 25-26 26-27 43-44 43-
48
44-45 45-46 46-47 47-48 60-61 60-65 61-62 62-63 63-64 64-65 73-74 73-75
74-75 81-82
81-86 82-83 83-84 84-85 85-86 126-127 126-128 127-128
exact/norm bonds :
1-20 4-8 5-35 9-153 9-154 15-156 22-30 25-29 27-41 43-54 60-70 66-67
73-74 73-75 74-75 81-88 106-107 110-113 115-150 116-117 117-118 117-120
126-127 126-128
127-128 135-136 136-137 138-139 139-140 143-144 143-145 143-146
exact bonds :
2-32 3-31 6-33 23-39 24-37 26-38 44-50 45-49 46-52 47-53 48-51 54-55
54-56 55-58 56-57 61-76 62-69 63-66 64-78 65-77 66-68 70-71 70-72 72-73
79-84 82-94 83-95
85-87 86-93 88-90 88-89 89-92 90-91 100-101 101-103 102-103 102-104 106-
108 109-110
110-111 111-112 121-122 121-123 124-125
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6 22-23 22-27 23-24 24-25 25-26 26-27 43-44 43-
48
44-45 45-46 46-47 47-48 60-61 60-65 61-62 62-63 63-64 64-65 81-82 81-86
82-83 83-84
84-85 85-86
isolated ring systems :
containing 1 : 22 : 43 : 60 : 81 :
G1:CN, NO2
G3:[*1],[*2]
G4:CF3,C1,NO2,CH3,OH,CN
G5:CH3,CF3,NO2,C1
G6:[*3],[*4],[*5],[*6],[*7]
G7:CH3,MeO,t-BuO
G8:CF3,OH,MeO,EtO,NH2,[*8],[*9],[*10],[*11],[*12],[*13],[*14],[*15]
G9:[*16],[*17],[*18],[*19],[*20]
Hydrogen count :
9:= exact 0 15:= exact 1
```

10/560012 9:3 E exact RC ring/chain 10:1 E exact RC ring/chain 11:1 E exact RC ring/chain 15:2 E exact RC ring/chain 79:1 E exact RC ring/chain 115:2 E exact RC ring/chain 125:2 E exact RC ring/chain 126:2 E exact RC ring/chain 128:2 E exact RC ring/chain 138:2 E exact RC ring/chain Match level : 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 8:CLASS 9:CLASS 10:CLASS 11:Atom 15:CLASS 20:CLASS 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:Atom 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS 35:CLASS 37:CLASS 38:CLASS 39:CLASS 41:CLASS 43:Atom 44:Atom 45:Atom 46:Atom 47:Atom 48:Atom 49:CLASS 50:CLASS 51:CLASS 52:CLASS 53:CLASS 54:CLASS 55:CLASS 56:CLASS 57:CLASS 58:CLASS 60:Atom 61:Atom 62:Atom 63:Atom 64:Atom 65:Atom 66:CLASS 67:CLASS 68:CLASS 69:CLASS 70:CLASS 71:CLASS 72:CLASS 73:Atom 74:Atom 75:Atom 76:CLASS 77:CLASS 78:CLASS 79:Atom 81:Atom 82:Atom 83:Atom 84:Atom 85:Atom 86:Atom 87:CLASS 88:CLASS 89:CLASS 90:CLASS 91:CLASS 92:CLASS 93:CLASS 94:CLASS 95:CLASS 99:CLASS 100:CLASS 101:CLASS 102:CLASS 103:CLASS 104:CLASS 106:CLASS 107:CLASS 108:CLASS 109:CLASS 110:CLASS 111:CLASS 112:CLASS 113:CLASS 115:CLASS 116:CLASS 117:CLASS 118:CLASS 120:CLASS 121:CLASS 122:CLASS 123:CLASS 124:CLASS 125:CLASS 126:Atom 127:Atom 128:Atom 135:CLASS 136:CLASS 137:CLASS 138:CLASS 139:CLASS 140:CLASS 143:CLASS 144:CLASS 145:CLASS 146:CLASS 150:CLASS 153:CLASS 154:CLASS 156:CLASS Generic attributes : 11: Saturation : Saturated 79: Saturation : Unsaturated Number of Carbon Atoms : less than 7 Type of Ring System : Monocyclic Element Count : Node 10: Limited C, C7 Node 11: Limited C.C6

Node 79: Limited

C,C6

Node 115: Limited C.C6

=> file zcaplus FILE 'ZCAPLUS' ENTERED AT 11:02:04 ON 12 MAR 2009 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.

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FILE COVERS 1907 - 12 Mar 2009 VOL 150 ISS 11 FILE LAST UPDATED: 11 Mar 2009 (20090311/ED)

ZCAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'ZCAPLUS' FILE

=> d stat que L69

151 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON (101130-93-2/BI OR 115416-50-7/BI OR 151951-35-8/BI OR 49645-18-3/BI OR 51332-25-3 /BI OR 821776-43-6/BI OR 821776-44-7/BI OR 821776-45-8/BI OR 821776-46-9/BI OR 821776-47-0/BI OR 821776-48-1/BI OR 821776-49 -2/BI OR 821776-50-5/BI OR 821776-51-6/BI OR 821776-52-7/BI OR 821776-53-8/BI OR 821776-54-9/BI OR 821776-55-0/BI OR 821776-56 -1/BI OR 821776-57-2/BI OR 821776-58-3/BI OR 821776-59-4/BI OR 821776-60-7/BI OR 821776-61-8/BI OR 821776-62-9/BI OR 821776-63 -0/BI OR 821776-64-1/BI OR 821776-65-2/BI OR 821776-66-3/BI OR 821776-67-4/BI OR 821776-68-5/BI OR 821776-69-6/BI OR 821776-70 -9/BI OR 821776-71-0/BI OR 821776-72-1/BI OR 821776-73-2/BI OR 821776-74-3/BI OR 821776-75-4/BI OR 821776-76-5/BI OR 821776-77 -6/BI OR 821776-78-7/BI OR 821776-79-8/BI OR 821776-80-1/BI OR 821776-81-2/BI OR 821776-82-3/BI OR 821776-83-4/BI OR 821776-84 -5/BI OR 821776-85-6/BI OR 821776-86-7/BI OR 821776-87-8/BI OR 821776-88-9/BI OR 821776-89-0/BI OR 821776-90-3/BI OR 821776-91 -4/BI OR 821776-92-5/BI OR 821776-93-6/BI OR 821776-94-7/BI OR 821776-95-8/BI OR 821776-96-9/BI OR 821776-97-0/BI OR 821776-98 -1/BI OR 821776-99-2/BI OR 821777-00-8/BI OR 821777-01-9/BI OR 821777-02-0/BI OR 821777-03-1/BI OR 821777-04-2/BI OR 821777-05 -3/BI OR 821777-06-4/BI OR 821777-07-5/BI OR 821777-08-6/BI OR 821777-09-7/BI OR 821777-10-0/BI OR 821777-11-1/BI OR 821777-12 -2/BI OR 821777-13-3/BI OR 821777-14-4/BI OR 821777-15-5/BI OR 821777-16-6/BI OR 821777-17-7/BI OR 821777-18-8/BI OR 821777-19 -9/BI OR 821777-20-2/BI OR 821777-21-3/BI OR 821777-22-4/BI OR 821777-23-5/BI OR 821777-24-6/BI OR 821777-25-7/BI OR 821777-26 -8/BI OR 821777-27-9/BI OR 821777-28-0/BI OR 821777-29-1/BI OR 821777-30-4/BI OR 821777-31-5/BI OR 821777-32-6/BI OR 821777-33 -7/BI OR 821777-34-8/BI OR 821777-35-9/BI OR 821777-36-0/BI OR 82177

I.14 SCR 616 L26 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

L27 SCR 1993

L32 SCR 1840 OR 2043 OR 1951

898352 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON (N>1 AND ((46.150.18/ L35 RID AND 1/NRS) OR (46.150.18/RID AND (1.13.1/RID OR 16.127.1/RI

D OR 46.150.1/RID) AND 2/NRS))) NOT PMS/CI L38 1054 SEA FILE=REGISTRY SUB=L35 SSS FUL L26 AND ((L14 AND L27) NOT

L32)

1.47 135 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L38 AND L3

16 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L3 NOT L47 L48 L50 1 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L48 AND BR/ELS AND

9/F

10 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L48 NOT BR/ELS L51 6 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L51 AND F/ELS L52 7 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L50 OR L52 L53

L54 142 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L47 OR L53

L55 65815 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON (C14H19N3O2/MF OR

C13H17N3O2/MF OR C12H13F3N2/MF OR C13H12F6N2O/MF OR C13H13N3O2/ MF OR C15H19F3N2/MF OR C11H11F3N2O2/MF OR C11H8F6N2/MF OR C12H11F3N2/MF OR C13H15F3N2/MF OR C13H15N3O2/MF OR C13H16N2O2/M

F OR C13H17CLN2/MF OR C13H17N3O3/MF OR C13H20N2O2/MF OR C14H14F6N2/MF OR C14H15F3N2O/MF OR C14H17F3N2/MF OR C14H17F3N2O 2/MF OR C14H17N3O2/MF OR C15H17F3N2/MF OR C15H17F3N2O/MF OR C15H17N3O2/MF OR C15H19F3N2O/MF OR C15H19F3N2O2/MF OR C16H17F3N 2/MF OR C16H21F3N2/MF OR C10H11F3N2O2/MF OR C10H11F3N2O3/MF OR

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C11H16N2O3/MF OR C11H7BRF9N/MF OR C12H10F6N2/MF OR C12H10F6N2O/ MF OR C12H11F5N2/MF OR C12H11F6N3/MF OR C12H13F3N2O/MF OR C12H16CLN3/MF OR C12H18CLN3O2/MF OR C12H18N2O3/MF OR C12H7F9N2/ MF OR C12H8F8N2/MF OR C12H9F6N5/MF OR C12H9F7N2/MF OR C13H12F6N

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C13H15CLN2/MF OR C13H15F3N2O2/MF OR C13H15F3N2O3/MF OR C13H17CLN2O/MF OR C13H17F3N2O2/MF OR C13H17F3N2O3/MF OR C13H17F3N2O4/MF OR C13H18N2O3/MF OR C13H20N2O3/MF OR C13H8F6N2/

MF OR C14H11F9N2O/MF OR C14H12F6N2/MF OR C14H13F3N2/MF OR C14H13F6N3O/MF OR C14H13F6N3O2/MF OR C14H14F6N2O/MF OR C14H15F3N2/MF OR C14H17CLN2/MF OR C14H17F3N2O/MF OR C14H17F3N2O

3/MF OR C14H17N3/MF OR C14H19CLN2/MF OR C14H19F3N2O2/MF OR C14H20N2O2/MF OR C14H22N2O2/MF OR C15H14F6N2/MF OR C15H16F6N2/M F OR C15H17F3N2O2/MF OR C15H19N3O2/MF OR C15H21CLN2/MF OR

C15H21CLN2O2/MF OR C15H21F3N2O2/MF OR C15H21N3O2/MF OR C15H24N2O2/MF OR C16H19F3N2/MF OR C16H19F3N2O2/MF OR C16H20F3NO /MF OR C16H21F3N2O/MF OR C16H21F3N2O2/MF OR C17H12F9N/MF OR

C17H19F6N3O2/MF OR C17H23F3N2/MF OR C17H25N3O2/MF OR C18H27F3N4 /MF OR C21H31F3N2OSI/MF OR C9H6F6N2O2/MF)

359 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L55 AND L38 L56 1.66 142 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L55 AND L54 366 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON L56 OR L66 L67

68 SEA FILE-ZCAPLUS SPE-ON ABB-ON PLU-ON L67 L68

L69 1 SEA FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L68 AND L1

=> d ibib abs hitstr L69 1

L69 ANSWER 1 OF 1 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2005:14359 ZCAPLUS Full-text

DOCUMENT NUMBER: 142:113710

TITLE: Preparation of substituted anilines as androgen

receptor modulators

INVENTOR(S): Blanc, Jean-Baptiste E.; Cadilla, Rodolfo; Cowan,
David John; Kaldor, Istvan; Larkin, Andrew L.;

Stewart, Eugene Lee; Turnbull, Philip Stewart; Trump,

Ryan Paul

PATENT ASSIGNEE(S): SmithKline Beecham Corporation, USA

SOURCE: PCT Int. Appl., 86 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.							DATE		APPLICATION NO.								
WO					A2				WO 2004-US18252								
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KΡ,	KR,	ΚZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	NI,
								PT,									
								UA,									
	RW:							ΜZ,									
								ТJ,									
		EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,	ΙT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
					BF,	ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,
			TD,														
EP								EP 2004-776383									
	R:							FR,									PT,
		ΙE,	SI,					CY,									
								JP 2006-533629									
US	US 20060148893				A1 20060706												
PRIORITY APPLN. INFO.:										003-							
										WO 2	004-	US18:	252	1	W 2	0040	609
OTHER SOURCE(S): GI					MARPAT 142:113710												

R4 R5 R6 R7

AB

This invention relates to non-steroidal compds. I [R1, R2, R4, R5 = H, CN, N02, halo, etc. (at least one of R1, R2, R4, R5 is not H); R3 = CN, N02, halo, etc.; R6, R7 = H, (Ra)xR9 (Ra = alkylene; x = 0-1; R9 = alkyl, haloalkyl,

hydroxyalkyl, etc.)] that are or are believed to be modulators of androgen, glucocorticoid, mineralocorticoid, and progesterone receptors, and also to the methods for the making and use of such compds. Thus, reacting 4-fluoro-2-trifluoromethylbenzonitrile with N-(cyclopropylmethyl)-N-propylamingfload 81% I [R1, R2, R5 = H; R3 = CN; R4 = CF3; R6 = CH2(cyclopropyl); R7 = Pr]. The compds. I are claimed to be useful in the treatment or prophylaxis of conditions or disorders that respond to selective androgen receptor modulation (no data given). The pharmaceutical composition comprising the compound I is disclosed.

- IT 821777-24-6P 821777-27-9P 821777-32-6P 821777-34-6P 821777-36-DE 821777-36-DE 821777-37-IP 821777-38-2P 821777-36-DE 821777-38-2P 821777-46-DE 821777-50-8P 821777-58-6P 821777-58-6P 821777-58-6P 821777-58-PE 821777-73-5P RI: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (USES) (preparation of substituted anilines as androgen receptor modulators)
- RN 821777-24-6 ZCAPLUS
 CN Benzonitrile, 4-[(cyclopropylmethyl)amino]-2-(trifluoromethyl)- (CA INDEX

- RN 821777-27-9 ZCAPLUS
- CN Benzenamine, N-(cyclopropylmethyl)-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-32-6 ZCAPLUS
- CN Benzonitrile, 4-[(1-cyclopropylethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-34-8 ZCAPLUS
- CN Benzonitrile, 4-[(2,2-dimethylpropyl)amino]-2-(trifluoromethyl)- (CA

CN

INDEX NAME)

RN 821777-36-0 ZCAPLUS

Benzonitrile, 4-[(2,2-dimethylpropyl)-2-propen-1-ylamino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-37-1 ZCAPLUS

CN Benzonitrile, 4-[(2,3-dihydroxypropyl)(2,2-dimethylpropyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-38-2 ZCAPLUS

CN Benzonitrile, 4-[(2,2-dimethylpropyl)(2-oxoethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-41-7 ZCAPLUS
- CN Benzonitrile, 4-[(1,1-dimethylethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-42-8 ZCAPLUS
- CN Benzenamine, N-(1,1-dimethylethyl)-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-44-0 ZCAPLUS
- CN Benzonitrile, 4-[(1,1-dimethylethyl)-2-propen-1-ylamino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-50-8 ZCAPLUS

- 821777-58-6 ZCAPLUS
- CN Benzonitrile, 4-[(2-hydroxyethyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl) - (CA INDEX NAME)

- 821777-66-6 ZCAPLUS RN
- Benzonitrile, 4-[(2-methyl-2-propen-1-yl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl) - (CA INDEX NAME)

- RN 821777-68-8 ZCAPLUS
- CN Benzonitrile, 4-[(3-methyl-2-buten-1-yl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl) - (CA INDEX NAME)

- 821777-73-5 ZCAPLUS
- CN Benzonitrile, 4-[(2-azidoethyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl) - (CA INDEX NAME)

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51332-25-3P 115416-50-7P 151951-35-8P
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821776-52-7P 821776-53-8P 821776-54-9P
821776-55-0P 821776-56-1P 821776-57-2P
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821777-35-9P 821777-39-3P 821777-40-6P
821777-43-9P 821777-45-1P 821777-46-2P
821777-47-3P 821777-48-4P 821777-49-5P
821777-51-9P 821777-52-0P 821777-53-1P
821777-55-3P 821777-56-4P 821777-57-5P
821777-59-7P 821777-60-0P 821777-61-1P
821777-62-2P 821777-63-3P 821777-64-4P
821777-65-5P 821777-67-7P 821777-69-9P
821777-70-2P 821777-71-3P 821777-72-4P
821777-74-6P 821777-75-7P 821777-76-8P
821777-77-9P 821777-78-0P 821778-33-0P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
```

(Uses)
(preparation of substituted anilines as androgen receptor modulators)

RN 51332-25-3 ZCAPLUS
CN Benzonitrile, 4-(dimethylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

N Benzoniciile, 4-(dimechylamino)-z-(criffuoromechyl)- (CA INDEX NAME)

RN 115416-50-7 ZCAPLUS

CN 1-Propanol, 3-[[4-nitro-3-(trifluoromethyl)phenyl]amino]- (CA INDEX NAME)

RN 151951-35-8 ZCAPLUS

CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

RN 821776-43-6 ZCAPLUS

CN Benzonitrile, 4-[(cyclopropylmethyl)propylamino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-44-7 ZCAPLUS

CN Benzonitrile, 4-[(cyclopropylmethyl)propylamino]-2-nitro- (CA INDEX NAME)

- RN 821776-45-8 ZCAPLUS
- CN Benzonitrile, 4-(di-2-propen-1-ylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

- n2cem cn— cn2
- RN 821776-46-9 ZCAPLUS
- CN Benzonitrile, 5-(cyclopentyl-2-propen-1-ylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-47-0 ZCAPLUS
- CN Benzonitrile, 4-(butylpropylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-48-1 ZCAPLUS
- CN Benzonitrile, 4-[ethyl(2-methyl-2-propen-1-yl)amino]-2-nitro- (CA INDEX NAME)

$$\begin{array}{c} \text{NO2} \\ \text{CN} \\ \text{Me} = \text{C} - \text{CH}_2 - \text{N} \\ \text{CH}_2 \\ \text{Et} \end{array}$$

821776-49-2 ZCAPLUS

CN Benzonitrile, 4-(butylethylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-50-5 ZCAPLUS

CN Benzonitrile, 4-(dipropylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-51-6 ZCAPLUS

CN Benzenamine, N-butyl-N-ethyl-3-methyl-4-nitro- (CA INDEX NAME)

RN 821776-52-7 ZCAPLUS

CN Benzonitrile, 5-(di-2-propen-1-ylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-53-8 ZCAPLUS
- CN Benzonitrile, 5-[(cyclopropylmethyl)propylamino]-2-nitro- (CA INDEX NAME)

- RN 821776-54-9 ZCAPLUS
- CN Benzonitrile, 4-(di-2-propen-1-ylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-55-0 ZCAPLUS
- CN Benzenamine, 3-methyl-4-nitro-N, N-dipropyl- (CA INDEX NAME)

- RN 821776-56-1 ZCAPLUS
- CN Benzonitrile, 4-[(1-methylpropyl)propylamino]-2-nitro- (CA INDEX NAME)

- RN 821776-57-2 ZCAPLUS
- CN Benzonitrile, 5-(butylethylamino)-2-nitro- (CA INDEX NAME)

RN 821776-58-3 ZCAPLUS

CN Benzonitrile, 2-chloro-4-[(cyclopropylmethyl)propylamino]- (CA INDEX NAME)

RN 821776-59-4 ZCAPLUS

CN Benzonitrile, 5-(butylpropylamino)-2-nitro- (CA INDEX NAME)

RN 821776-60-7 ZCAPLUS

CN Benzonitrile, 5-[(2-methoxyethyl)methylamino]-2-nitro- (CA INDEX NAME)

RN 821776-61-8 ZCAPLUS

CN Benzonitrile, 2-chloro-4-(di-2-propen-1-ylamino)- (CA INDEX NAME)

RN 821776-62-9 ZCAPLUS

CN Benzonitrile, 4-[(2-methoxyethyl)propylamino]-2-nitro- (CA INDEX NAME)

RN 821776-63-0 ZCAPLUS

CN Benzonitrile, 4-(cyclopenty1-2-propen-1-ylamino)-2-nitro- (CA INDEX NAME)

RN 821776-64-1 ZCAPLUS

RN 821776-65-2 ZCAPLUS

CN Benzonitrile, 2-chloro-4-[[2-(dimethylamino)ethyl]methylamino]- (CA INDEX NAME)

- RN 821776-66-3 ZCAPLUS
- CN Benzenamine, N-(2-methoxyethyl)-N,2-dimethyl-4-nitro- (CA INDEX NAME)

- RN 821776-67-4 ZCAPLUS
- CN Benzenamine, N-cyclopentyl-4-nitro-N-2-propen-1-yl-3-(trifluoromethyl)-(CA INDEX NAME)

- RN 821776-68-5 ZCAPLUS
- CN Phenol, 5-(dipropylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-69-6 ZCAPLUS
- CN Benzonitrile, 2-chloro-4-(dipropylamino)- (CA INDEX NAME)

RN 821776-70-9 ZCAPLUS

Benzonitrile, 5-[ethyl(2-methyl-2-propen-1-yl)amino]-2-nitro- (CA INDEX NAME)

RN 821776-71-0 ZCAPLUS

CN Benzenamine, 4-nitro-N, N-di-2-propen-1-yl-3-(trifluoromethyl)- (CA INDEX NAME)

821776-72-1 ZCAPLUS RN

CN Benzenamine, 2-methyl-4-nitro-N, N-di-2-propen-1-yl- (CA INDEX NAME)

821776-73-2 ZCAPLUS RN

CN Phenol, 5-[(cyclopropylmethyl)propylamino]-2-nitro- (CA INDEX NAME)

RN 821776-74-3 ZCAPLUS

CN Benzenamine, N-(2-methoxyethyl)-3-methyl-4-nitro-N-propyl- (CA INDEX NAME)

RN 821776-75-4 ZCAPLUS

CN Benzonitrile, 4-(butylpropylamino)-2-chloro- (CA INDEX NAME)

RN 821776-76-5 ZCAPLUS

CN Benzenamine, N-buty1-2-chloro-N-methy1-4-nitro- (CA INDEX NAME)

RN 821776-77-6 ZCAPLUS

CN Benzonitrile, 4-(dipropylamino)-2-nitro- (CA INDEX NAME)

RN 821776-78-7 ZCAPLUS

CN Benzonitrile, 2-chloro-4-[(2-methoxyethyl)propylamino]- (CA INDEX NAME)

RN 821776-79-8 ZCAPLUS

CN Benzenamine, 3-methyl-4-nitro-N,N-di-2-propen-1-yl- (CA INDEX NAME)

RN 821776-80-1 ZCAPLUS

CN Benzenamine, N-(1-methylpropyl)-4-nitro-N-propyl-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-81-2 ZCAPLUS

CN Benzonitrile, 4-(dipentylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-82-3 ZCAPLUS
- CN Benzenamine, N-(cyclopropylmethyl)-3-methyl-4-nitro-N-propyl- (CA INDEX NAME)

- RN 821776-83-4 ZCAPLUS
- CN Benzonitrile, 5-(dipropylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-84-5 ZCAPLUS
- CN Benzenamine, N, N-dibutyl-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N2,N2-trimethyl- (CA INDEX NAME)

RN 821776-86-7 ZCAPLUS

CN Benzonitrile, 5-[(1-methylpropyl)propylamino]-2-nitro- (CA INDEX NAME)

RN 821776-87-8 ZCAPLUS

CN Benzenamine, N-ethyl-N-(2-methyl-2-propen-1-yl)-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-88-9 ZCAPLUS

CN Benzenamine, N-(2-methoxyethyl)-4-nitro-N-propyl-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-89-0 ZCAPLUS

CN Benzonitrile, 4-(butylethylamino)-2-chloro- (CA INDEX NAME)

- RN 821776-90-3 ZCAPLUS
- CN Benzonitrile, 5-(dibutylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-91-4 ZCAPLUS
- CN Benzonitrile, 4-(butylethylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-92-5 ZCAPLUS
- CN Benzonitrile, 4-[(2-methoxyethy1)propylamino]-2-(trifluoromethy1)- (CA INDEX NAME)

- меошсидшена
- RN 821776-93-6 ZCAPLUS
- CN Benzonitrile, 5-[(2-methoxyethy1)propylamino]-2-nitro- (CA INDEX NAME)

RN 821776-94-7 ZCAPLUS

CN Benzonitrile, 4-(cyclopenty1-2-propen-1-ylamino)-2-(trifluoromethy1)- (CA INDEX NAME)

RN 821776-95-8 ZCAPLUS

CN Benzonitrile, 4-(butylpropylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821776-96-9 ZCAPLUS

CN Benzenamine, N, N-dibutyl-3-methyl-4-nitro- (CA INDEX NAME)

RN 821776-97-0 ZCAPLUS

CN Benzonitrile, 4-(hexylmethylamino)-2-nitro- (CA INDEX NAME)

- RN 821776-98-1 ZCAPLUS
- CN Benzonitrile, 4-(dibutylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821776-99-2 ZCAPLUS
- CN Benzenamine, N-(cyclopropylmethyl)-4-nitro-N-propyl-2-(trifluoromethyl)-(CA INDEX NAME)

- RN 821777-00-8 ZCAPLUS
- CN Benzenamine, N-cyclohexyl-4-nitro-N-2-propen-1-yl-3-(trifluoromethyl)-(CA INDEX NAME)

- RN 821777-01-9 ZCAPLUS
- CN Benzonitrile, 4-[(2-methoxyethyl)methylamino]-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-02-0 ZCAPLUS

CN Benzonitrile, 4-(di-2-propen-1-ylamino)-3-nitro- (CA INDEX NAME)

RN 821777-03-1 ZCAPLUS

CN 1,3-Propanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N3,N3-trimethyl- (CA INDEX NAME)

RN 821777-04-2 ZCAPLUS

CN Benzenamine, N,N-bis(2-methoxyethyl)-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-05-3 ZCAPLUS

CN Benzonitrile, 4-[(1-methylpropyl)propylamino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-06-4 ZCAPLUS

CN Benzonitrile, 2-chloro-4-[ethyl(2-methyl-2-propen-1-yl)amino]- (CA INDEX NAME)

RN 821777-07-5 ZCAPLUS

CN Benzenamine, N-cyclohexyl-N-ethyl-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-08-6 ZCAPLUS

CN Benzonitrile, 2-chloro-4-(dibutylamino)- (CA INDEX NAME)

RN 821777-09-7 ZCAPLUS

CN Benzonitrile, 4-(cyclohexylethylamino)-2-nitro- (CA INDEX NAME)

RN 821777-10-0 ZCAPLUS

CN Benzonitrile, 4-[bis(2-ethoxyethyl)amino]-3-chloro- (CA INDEX NAME)

RN 821777-11-1 ZCAPLUS

CN Benzenamine, N-butvl-N-ethvl-4-nitro-3-(trifluoromethvl)- (CA INDEX NAME)

RN 821777-12-2 ZCAPLUS

CN Benzenamine, 3-methyl-N-(1-methylpropyl)-4-nitro-N-propyl- (CA INDEX NAME)

RN 821777-13-3 ZCAPLUS

CN Benzenamine, N-(2-methoxyethyl)-N-methyl-4-nitro-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-15-5 ZCAPLUS

CN Benzonitrile, 4-[bis[3-(dimethylamino)propyl]amino]-2-(trifluoromethyl)-(CA INDEX NAME)

RN 821777-16-6 ZCAPLUS

CN Benzenamine, N-(cyclopropylmethyl)-4-nitro-N-propyl-3-(trifluoromethyl)-(CA INDEX NAME)

RN 821777-17-7 ZCAPLUS

CN Benzonitrile, 4-(methyloctylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-18-8 ZCAPLUS

CN Benzonitrile, 4-(propylamino)-2-(trifluoromethy1)- (CA INDEX NAME)

RN 821777-19-9 ZCAPLUS

CN Benzenamine, 4-nitro-N-propyl-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-20-2 ZCAPLUS

CN Benzonitrile, 4-[(3-hydroxypropyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-21-3 ZCAPLUS

CN Benzonitrile, 4-[[2-hydroxy-1-(hydroxymethyl)ethyl]amino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-22-4 ZCAPLUS

CN Benzonitrile, 4-(diethylamino)-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-23-5 ZCAPLUS
- CN Benzonitrile, 4-[methyl(2-methylpropyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-25-7 ZCAPLUS
- CN Benzonitrile, 4-[(cyclopropylmethyl)(3-hydroxypropyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-26-8 ZCAPLUS
- CN Benzonitrile, 4-[(cyclopropylmethyl)(2-hydroxyethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-28-0 ZCAPLUS
- CN 1-Propanol, 3-[(cyclopropylmethyl)[4-nitro-3-(trifluoromethyl)phenyl]amino]- (CA INDEX NAME)

- RN 821777-29-1 ZCAPLUS
 - Ethanol, 2-[(cyclopropylmethyl)[4-nitro-3-(trifluoromethyl)phenyl]amino]-(CA INDEX NAME)

- RN 821777-30-4 ZCAPLUS
- CN Benzonitrile, 4-[(cyclopropylmethyl)amino]-3-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-31-5 ZCAPLUS

- RN 821777-33-7 ZCAPLUS
- CN Benzonitrile, 4-[(1-cyclopropylethy1)-2-propen-1-ylamino]-2-(trifluoromethy1)- (CA INDEX NAME)

- RN 821777-35-9 ZCAPLUS

- RN 821777-39-3 ZCAPLUS
- CN Benzonitrile, 4-[(2,2-dimethylpropyl)(2-hydroxyethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-40-6 ZCAPLUS
- CN Benzonitrile, 4-[(2,2-dimethylpropyl)propylamino]-2-(trifluoromethyl)-(CA INDEX NAME)

- RN 821777-43-9 ZCAPLUS
- CN Benzenamine, N-(cyclopropylmethyl)-N-(1,1-dimethylethyl)-4-nitro-3-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-45-1 ZCAPLUS

RN 821777-46-2 ZCAPLUS

CN Benzonitrile, 4-[(1,1-dimethylethyl)propylamino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-47-3 ZCAPLUS

CN Benzonitrile, 4-[(3-hydroxypropyl)](1S)-1-methylpropyl]amino]-2-(trifluoromethyl)- (CA INDEX NAME)

Absolute stereochemistry.

- RN 821777-48-4 ZCAPLUS
- CN Benzonitrile, 2-(trifluoromethyl)-4-[(3,3,3-trifluoropropyl)amino]- (CA INDEX NAME)

F3C_ CH2_ CH2

- 821777-49-5 ZCAPLUS RN
- CN Benzonitrile, 4-[bis(2-fluoroethy1)amino]-2-(trifluoromethy1)- (CA INDEX NAME)

- RN 821777-51-9 ZCAPLUS
- CN Benzenamine, 4-nitro-N-(2,2,2-trifluoroethyl)-3-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-52-0 ZCAPLUS
- CN Benzonitrile, 4-[(3-hydroxypropyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl) - (CA INDEX NAME)

- RN 821777-53-1 ZCAPLUS
- CN Benzonitrile, 4-[(3-hydroxypropyl)(2,2,2-trifluoroethyl)amino]-3-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-55-3 ZCAPLUS
- CN Benzonitrile, 4-[bis(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-56-4 ZCAPLUS
- CN Benzonitrile, 4-[(2,2-difluoroethyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-57-5 ZCAPLUS
- CN [1,1'-Biphenyl]-4-amine, N,N-bis(2,2,2-trifluoroethyl)-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-59-7 ZCAPLUS

CN Benzonitrile, 4-[(2-methoxyethyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-60-0 ZCAPLUS

CN Benzonitrile, 4-[(2-ethoxyethy1)(2,2,2-trifluoroethy1)amino]-2-(trifluoromethy1)- (CA INDEX NAME)

RN 821777-61-1 ZCAPLUS

CN Benzonitrile, 4-[[2-(2,2,2-trifluoroethoxy)ethy1](2,2,2-trifluoroethy1)amino]-2-(trifluoromethy1)- (CA INDEX NAME)

- RN 821777-62-2 ZCAPLUS
- CN Benzonitrile, 4-[methyl(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)-(CA INDEX NAME)

- RN 821777-63-3 ZCAPLUS

- RN 821777-64-4 ZCAPLUS
- CN Benzonitrile, 4-[propyl(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)-(CA INDEX NAME)

- RN 821777-65-5 ZCAPLUS

RN 821777-67-7 ZCAPLUS

CN Benzonitrile, 4-[(2-methylpropyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-69-9 ZCAPLUS

CN Benzonitrile, 4-[(3-methylbutyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

RN 821777-70-2 ZCAPLUS

CN Benzonitrile, 4-[2-propyn-1-yl(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-71-3 ZCAPLUS
- CN Benzonitrile, 4-[(2-fluoroethyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-72-4 ZCAPLUS
- CN Benzonitrile, 4-[[2-(methylthio)ethyl](2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-74-6 ZCAPLUS
- CN Acetamide, N-[2-[[4-cyano-3-(trifluoromethyl)phenyl](2,2,2-trifluoroethyl)amino]ethyl]- (CA INDEX NAME)

- RN 821777-75-7 ZCAPLUS
- CN Carbamic acid, [2-[[4-cyano-3-(trifluoromethyl)phenyl](2,2,2-trifluoroethyl)amino]ethyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 821777-76-8 ZCAPLUS

CN Carbamic acid, [2-[[4-cyano-3-(trifluoromethyl)phenyl](2,2,2-trifluoroethyl)amino]ethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 821777-77-9 ZCAPLUS

CN Methanesulfonamide, N-[2-[[4-cyano-3-(trifluoromethyl)phenyl]](2,2,2trifluoroethyl)amino]ethyl]- (CA INDEX NAME)

RN 821777-78-0 ZCAPLUS

CN 1,2-Benzenedicarbonitrile, 4-(dipropylamino)- (CA INDEX NAME)

RN 821778-33-0 ZCAPLUS

CN Benzenamine, 4-bromo-N, N-bis(2,2,2-trifluoroethyl)-3-(trifluoromethyl)-(CA INDEX NAME)

IT 821777-79-1P 821777-80-4P 821777-85-9P

821777-86-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of substituted anilines as androgen receptor modulators) RN $\,$ 821777-79-1 $\,$ ZCAPLUS

CN Benzonitrile, 4-[(cyclopropylmethyl)[3-[[(1,1-

dimethylethyl)dimethylsilyl]oxy]propyl]amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-80-4 ZCAPLUS
- CN Benzonitrile, 4-[[(1S)-1-methylpropyl]amino]-2-(trifluoromethyl)- (CA INDEX NAME)

Absolute stereochemistry.

- RN 821777-85-9 ZCAPLUS
- CN Benzonitrile, 4-[[2-[(methylsulfonyl)oxy]ethyl](2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

- RN 821777-86-0 ZCAPLUS
- CN Benzonitrile, 4-[(2-aminoethyl)(2,2,2-trifluoroethyl)amino]-2-(trifluoromethyl)- (CA INDEX NAME)

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chain nodes :
8 9 10 11 15 20 29 30 31 32 33 35 37 38 39 41 49 50 51 52 53
54 55 56 57 58 66 67 68 69 70 71 72 76 77 78 79 87 88 89 90 91
92 93 94 95
99 100 101 102 103 104 106 107 108 109 110 111 112 113 115 116 117
118 120 121
122 123 124 125 135 136 137 138 139 140 143 144 145 146 150 153 154
156
ring nodes :
1 2 3 4 5 6 22 23 24 25 26 27 43 44 45 46 47 48 60 61 62 63
64 65 73 74 75 81 82 83 84 85 86 126 127 128
chain bonds :
1-20 2-32 3-31 4-8 5-35 6-33 9-153 9-154 15-156 22-30 23-39 24-37 25-29
26-38 27-41 43-54 44-50 45-49 46-52 47-53 48-51 54-55 54-56 55-58 56-57
60-70 61-76
62-69 63-66 64-78 65-77 66-67 66-68 70-71 70-72 72-73 79-84 81-88 82-94
83-95 85-87
86-93 88-90 88-89 89-92 90-91 100-101 101-103 102-103 102-104 106-108
106-107 109-110
110-111 110-113 111-112 115-150 116-117 117-118 117-120 121-122 121-123
124-125 135-136
136-137 138-139 139-140 143-144 143-145 143-146
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6 22-23 22-27 23-24 24-25 25-26 26-27 43-44 43-
48
44-45 45-46 46-47 47-48 60-61 60-65 61-62 62-63 63-64 64-65 73-74 73-75
74-75 81-82
81-86 82-83 83-84 84-85 85-86 126-127 126-128 127-128
exact/norm bonds :
1-20 4-8 5-35 9-153 9-154 15-156 22-30 25-29 27-41 43-54 60-70 66-67
73-74 73-75 74-75 81-88 106-107 110-113 115-150 116-117 117-118 117-120
126-127 126-128
127-128 135-136 136-137 138-139 139-140 143-144 143-145 143-146
exact bonds :
2-32 3-31 6-33 23-39 24-37 26-38 44-50 45-49 46-52 47-53 48-51 54-55
54-56 55-58 56-57 61-76 62-69 63-66 64-78 65-77 66-68 70-71 70-72 72-73
79-84 82-94 83-95
85-87 86-93 88-90 88-89 89-92 90-91 100-101 101-103 102-103 102-104 106-
108 109-110
110-111 111-112 121-122 121-123 124-125
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6 22-23 22-27 23-24 24-25 25-26 26-27 43-44 43-
48
44-45 45-46 46-47 47-48 60-61 60-65 61-62 62-63 63-64 64-65 81-82 81-86
82-83 83-84
84-85 85-86
isolated ring systems :
containing 1 : 22 : 43 : 60 : 81 :
```

G1:CN, NO2

G3:[*1],[*2]

G4:CF3,C1,NO2,CH3,OH,CN

G5:CH3,CF3,NO2,C1

G6:[*3],[*4],[*5],[*6],[*7]

```
G7:CH3,MeO,t-BuO
```

```
G8:CF3,OH,MeO,EtO,NH2,[*8],[*9],[*10],[*11],[*12],[*13],[*14],[*15]
```

G9:[*16],[*17],[*18],[*19],[*20]

Hydrogen count :

9:= exact 0 15:= exact 1

Connectivity:

9:3 E exact RC ring/chain 10:1 E exact RC ring/chain 11:1 E exact RC ring/chain 15:2 E exact RC ring/chain 79:1 E exact RC ring/chain 115:2 E exact RC ring/chain 125:2 E exact RC ring/chain 126:2 E exact RC ring/chain 128:2 E exact RC ring/chain 138:2 E

exact RC ring/chain

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 8:CLASS 9:CLASS 10:CLASS 11:Atom 15:CLASS 20:CLASS 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:Atom 29:CLASS 30:CLASS 31:CLASS

32:CLASS 33:CLASS 35:CLASS 37:CLASS 38:CLASS 39:CLASS 41:CLASS 43:Atom 44:Atom 45:Atom

46:Atom 47:Atom 48:Atom 49:CLASS 50:CLASS 51:CLASS 52:CLASS 53:CLASS 54:CLASS 55:CLASS

56:CLASS 57:CLASS 58:CLASS 60:Atom 61:Atom 62:Atom 63:Atom 64:Atom 65:Atom 66:CLASS

67:CLASS 68:CLASS 69:CLASS 70:CLASS 71:CLASS 72:CLASS 73:Atom 74:Atom 75:Atom 76:CLASS 77:CLASS

78:CLASS 79:Atom 81:Atom 82:Atom 83:Atom 84:Atom 85:Atom 86:Atom 87:CLASS 88:CLASS

89:CLASS 90:CLASS 91:CLASS 92:CLASS 93:CLASS 94:CLASS 95:CLASS 99:CLASS 100:CLASS 101:CLASS

102:CLASS 103:CLASS 104:CLASS 106:CLASS 107:CLASS 108:CLASS 109:CLASS 110:CLASS 111:CLASS

112:CLASS 113:CLASS 115:CLASS 116:CLASS 117:CLASS 118:CLASS 120:CLASS

121:CLASS 122:CLASS 123:CLASS 124:CLASS 125:CLASS 126:Atom 127:Atom 128:Atom 135:CLASS 136:CLASS

137:CLASS 138:CLASS 139:CLASS 140:CLASS 143:CLASS 144:CLASS 145:CLASS 146:CLASS

150:CLASS 153:CLASS 154:CLASS 156:CLASS

Generic attributes :

: Saturated Saturation

79:

Saturation : Unsaturated Number of Carbon Atoms : less than 7 Type of Ring System : Monocyclic

Element Count : Node 10: Limited C.C7

Node 11: Limited C.C6

Node 79: Limited C, C6

Node 115: Limited C. C6

L14

L26

=> d stat que L70

L3 151 SEA FILE=REGISTRY SPE=ON ABB=ON PLU=ON (101130-93-2/BI OR 115416-50-7/BI OR 151951-35-8/BI OR 49645-18-3/BI OR 51332-25-3 /BI OR 821776-43-6/BI OR 821776-44-7/BI OR 821776-45-8/BI OR 821776-46-9/BI OR 821776-47-0/BI OR 821776-48-1/BI OR 821776-49 -2/BI OR 821776-50-5/BI OR 821776-51-6/BI OR 821776-52-7/BI OR 821776-53-8/BT OR 821776-54-9/BT OR 821776-55-0/BT OR 821776-56 -1/BI OR 821776-57-2/BI OR 821776-58-3/BI OR 821776-59-4/BI OR 821776-60-7/BI OR 821776-61-8/BI OR 821776-62-9/BI OR 821776-63 -0/BI OR 821776-64-1/BI OR 821776-65-2/BI OR 821776-66-3/BI OR 821776-67-4/BI OR 821776-68-5/BI OR 821776-69-6/BI OR 821776-70 -9/BI OR 821776-71-0/BI OR 821776-72-1/BI OR 821776-73-2/BI OR 821776-74-3/BI OR 821776-75-4/BI OR 821776-76-5/BI OR 821776-77 -6/BI OR 821776-78-7/BI OR 821776-79-8/BI OR 821776-80-1/BI OR 821776-81-2/BI OR 821776-82-3/BI OR 821776-83-4/BI OR 821776-84 -5/BI OR 821776-85-6/BI OR 821776-86-7/BI OR 821776-87-8/BI OR 821776-88-9/BI OR 821776-89-0/BI OR 821776-90-3/BI OR 821776-91 -4/BI OR 821776-92-5/BI OR 821776-93-6/BI OR 821776-94-7/BI OR 821776-95-8/BI OR 821776-96-9/BI OR 821776-97-0/BI OR 821776-98 -1/BI OR 821776-99-2/BI OR 821777-00-8/BI OR 821777-01-9/BI OR 821777-02-0/BI OR 821777-03-1/BI OR 821777-04-2/BI OR 821777-05 -3/BI OR 821777-06-4/BI OR 821777-07-5/BI OR 821777-08-6/BI OR 821777-09-7/BI OR 821777-10-0/BI OR 821777-11-1/BI OR 821777-12 -2/BI OR 821777-13-3/BI OR 821777-14-4/BI OR 821777-15-5/BI OR 821777-16-6/BI OR 821777-17-7/BI OR 821777-18-8/BI OR 821777-19 -9/BI OR 821777-20-2/BI OR 821777-21-3/BI OR 821777-22-4/BI OR 821777-23-5/BI OR 821777-24-6/BI OR 821777-25-7/BI OR 821777-26 -8/BI OR 821777-27-9/BI OR 821777-28-0/BI OR 821777-29-1/BI OR 821777-30-4/BI OR 821777-31-5/BI OR 821777-32-6/BI OR 821777-33 -7/BI OR 821777-34-8/BI OR 821777-35-9/BI OR 821777-36-0/BI OR 82177

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation

SCR 616

STR

Stru	cture attri	butes	s must be viewe	ed using	STN Exp	ress que:	ry preparation.	
L27		SCR	1993					
L32		SCR	1840 OR 2043 (OR 1951				
L35	898352	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	(N>1 AND ((46.150.	18/
		RID	AND 1/NRS) OR	(46.150	.18/RID .	AND (1.1	3.1/RID OR 16.127.1	/RI
		D O	R 46.150.1/RII) AND 2	/NRS)))	NOT PMS/	CI	
L38	1054	SEA	FILE=REGISTRY	SUB=L35	SSS FUL	L26 AND	((L14 AND L27) NOT	
		L32)					
L47	135	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L38 AND L3	
L48	16	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L3 NOT L47	
L50	1	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L48 AND BR/ELS AND)
		9/F						
L51	10	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L48 NOT BR/ELS	
L52	6	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L51 AND F/ELS	
L53	7	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L50 OR L52	
L54	142	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	L47 OR L53	
L55	65815	SEA	FILE=REGISTRY	SPE=ON	ABB=ON	PLU=ON	(C14H19N3O2/MF OR	

C13H17N3O2/MF OR C12H13F3N2/MF OR C13H12F6N2O/MF OR C13H13N3O2/

MF OR C15H19F3N2/MF OR C11H11F3N2O2/MF OR C11H8F6N2/MF OR C12H11F3N2/MF OR C13H15F3N2/MF OR C13H15N3O2/MF OR C13H16N2O2/M F OR C13H17CLN2/MF OR C13H17N3O3/MF OR C13H20N2O2/MF OR C14H14F6N2/MF OR C14H15F3N2O/MF OR C14H17F3N2/MF OR C14H17F3N2O 2/MF OR C14H17N3O2/MF OR C15H17F3N2/MF OR C15H17F3N2O/MF OR C15H17N3O2/MF OR C15H19F3N2O/MF OR C15H19F3N2O2/MF OR C16H17F3N 2/MF OR C16H21F3N2/MF OR C10H11F3N2O2/MF OR C10H11F3N2O3/MF OR C10H13CLN2O3/MF OR C10H6F6N2/MF OR C10H9F3N2/MF OR C11H11F3N2/M F OR C11H11F3N2O/MF OR C11H13F3N2O2/MF OR C11H13F3N2O3/MF OR C11H13N3O3/MF OR C11H15CLN2O2/MF OR C11H16CLN3O2/MF OR C11H16N2O3/MF OR C11H7BRF9N/MF OR C12H10F6N2/MF OR C12H10F6N2O/ MF OR C12H11F5N2/MF OR C12H11F6N3/MF OR C12H13F3N2O/MF OR C12H16CLN3/MF OR C12H18CLN3O2/MF OR C12H18N2O3/MF OR C12H7F9N2/ MF OR C12H8F8N2/MF OR C12H9F6N5/MF OR C12H9F7N2/MF OR C13H12F6N 2/MF OR C13H12F6N2O3S/MF OR C13H12F6N2S/MF OR C13H13CLN2/MF OR C13H13F3N2/MF OR C13H13F3N2O2/MF OR C13H13F6N3O2S/MF OR C13H15CLN2/MF OR C13H15F3N2O2/MF OR C13H15F3N2O3/MF OR C13H17CLN2O/MF OR C13H17F3N2O2/MF OR C13H17F3N2O3/MF OR C13H17F3N2O4/MF OR C13H18N2O3/MF OR C13H20N2O3/MF OR C13H8F6N2/ MF OR C14H11F9N2O/MF OR C14H12F6N2/MF OR C14H13F3N2/MF OR C14H13F6N3O/MF OR C14H13F6N3O2/MF OR C14H14F6N2O/MF OR C14H15F3N2/MF OR C14H17CLN2/MF OR C14H17F3N2O/MF OR C14H17F3N2O 3/MF OR C14H17N3/MF OR C14H19CLN2/MF OR C14H19F3N2O2/MF OR C14H20N2O2/MF OR C14H22N2O2/MF OR C15H14F6N2/MF OR C15H16F6N2/M F OR C15H17F3N2O2/MF OR C15H19N3O2/MF OR C15H21CLN2/MF OR C15H21CLN2O2/MF OR C15H21F3N2O2/MF OR C15H21N3O2/MF OR C15H24N2O2/MF OR C16H19F3N2/MF OR C16H19F3N2O2/MF OR C16H20F3NO /MF OR C16H21F3N2O/MF OR C16H21F3N2O2/MF OR C17H12F9N/MF OR C17H19F6N3O2/MF OR C17H23F3N2/MF OR C17H25N3O2/MF OR C18H27F3N4

		/ MF	OR CZIH3IF3NZOSI/MF OR C9H6F6NZOZ/MF)
L56	359	SEA	FILE=REGISTRY SPE=ON ABB=ON PLU=ON L55 AND L38
L57	68	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L56
L58	49	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L57 AND P/DT
L59	19	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L57 NOT L58
L60	16	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L59 AND PY<2004
L61	28	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L58 AND PD<20030610
L62	35	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L58 AND PRD<20030610
L63	30	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L58 AND AD<20030610
L64	51	SEA	FILE-ZCAPLUS SPE=ON ABB=ON PLU=ON (L60 OR L61 OR L62 OR
		L63)
L66	142	SEA	FILE=REGISTRY SPE=ON ABB=ON PLU=ON L55 AND L54
L67	366	SEA	FILE=REGISTRY SPE=ON ABB=ON PLU=ON L56 OR L66
L68	68	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L67
L70	51	SEA	FILE=ZCAPLUS SPE=ON ABB=ON PLU=ON L68 AND L64

=> file zcaplus
FILE 'ZCAPLUS' ENTERED AT 11:02:50 ON 12 MAR 2009
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FILE COVERS 1907 - 12 Mar 2009 VOL 150 ISS 11 FILE LAST UPDATED: 11 Mar 2009 (20090311/ED)

ZCAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'ZCAPLUS' FILE

=> d ibib abs hitstr L70 1-51

L70 ANSWER 1 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2006:1342415 ZCAPLUS Full-text

DOCUMENT NUMBER: 146:89971

TITLE: Nile red luminescent compound emitting red light, process for producing the same and luminescence

element utilizing the same and luminescence

INVENTOR(S): Nakaya, Tadao; Tajima, Akio; Saikawa, Tomoyuki;

Takano, Shinji; Yamauchi, Takao; Mori, Hidemasa
PATENT ASSIGNEE(S): Hirose Engineering Co., Ltd, Japan

SOURCE: U.S. Pat. Appl. Publ., 55pp., Division of U.S. Ser.

No. 501,398.

CODEN: USXXCO

LANGUAGE: English
FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

DOCUMENT TYPE:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20060287524 US 20050113575	A1 A1	20061221 20050526	US 2006-486111 US 2004-501398	20060831 20040715 <
PRIORITY APPLN. INFO.:	AI	20030326	US 2004-501398 F	3 20040715
			JP 2002-12222 F JP 2002-12224 F	20020121 <
			JP 2002-14881 F	20020123 <
			JP 2002-172127 F	20020612 <
			WO 2003-JP477 V	20030121 <

OTHER SOURCE(S): MARPAT 146:89971

AB Luminescent derivs. of 9-amino-5H-benzo[a]phenoxazin-5-one (Nile Red derivs.) and methods for producing them are described. Electroluminescent devices employing the compds. are also described. The patent also describes the preparation of unclaimed luminescent derivs. of 9-amino-5H-benzo[a]benzo[a]benzo[hizzin-5-one]

IT 398482-49-0

RL: RCT (Reactant); RACT (Reactant or reagent) (luminescent aminobenzophenoxazinone derivs. and their preparation and light-emitting devices using them)

RN 398482-49-0 ZCAPLUS

CN Phenol, 5-[bis(1-methylethyl)amino]-2-nitro- (CA INDEX NAME)

L70 ANSWER 2 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2005:902613 ZCAPLUS Full-text

ACCESSION NUMBER: 2005:902613 20

DOCUMENT NUMBER: 143:248160

TITLE: Preparation of benzamides for promoting apoptosis
INVENTOR(S): Bajji, Ashok C.; Arranz, Esther; Srinlvassan, Jayasree
M.; Delmar, Eric; Slade, Rachel; Willardsen, Jon Adam

PATENT ASSIGNEE(S): Myriad Genetics, Incorporated, USA

SOURCE: Wyliad Generics, Incorporated, USA

Wyliad Generics, Incorporated, USA

U.S. Pat. Appl. Publ., 121 pp., Cont.-in-part of Appl.

No. PCT/US03/22183.

CODEN: USXXCO
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

GT

	PATENT NO.				KIND DATE						ION				ATE			
US WO	2005 2004 2004	0187 0068	300 58		A1 A2		2005	0825 0122		US 2	005-	3927 US22	5		2	0050	118 · 715 ·	
	W: RW:	CO, GM, LS, PG, TR, GH, KG,	CR, HR, LT, PH, TT, GM, KZ,	CU, HU, LU, PL, TZ, KE, MD,	CZ, ID, LV, PT, UA, LS, RU,	DE, IL, MA, RO, UG, MW, TJ,	DK, IN, MD, RU, US, MZ, TM,	AZ, DM, IS, MG, SC, UZ, SD, AT, IT,	DZ, JP, MK, SD, VC, SL, BE,	EC, KE, MN, SE, VN, SZ, BG,	EE, KG, MW, SG, YU, TZ, CH,	ES, KP, MX, SK, ZA, UG, CY,	FI, KR, MZ, SL, ZM, ZM, CZ,	GB, KZ, NI, SY, ZW, ZW, DE,	GD, LC, NO, TJ, AM, DK,	GE, LK, NZ, TM, AZ, EE,	GH, LR, OM, TN, BY, ES,	
BF, BJ, CF, PRIORITY APPLN. INFO.: OTHER SOURCE(S):					GR, HU, IE, IT, CG, CI, CM, GA,				US 2 US 2 WO 2	002- 002- 003-	3962 3967 US22	66P 73P 183		SN, P 2 P 2 A2 2	0020 0020	715 · 716 ·		

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Benzamides, e.g. I (R2, R3 = halo, C1-C6-haloalky1; R4, R5 = C1-C6-alky1; R6 = H, halo; R7 = halo, C1-C4-haloalky1) and II (R11, R12, R13, R14, R15, R16, R17, R18 = H, halo, N3, OH, SH, cyano, C1-C6-(halo) (hydroxy)alky1, C2-C6-alkeny1, C2-C6-alkeny1, C2-C6-alkeny1, C2-C6-alkeny1, C2-C6-alky1, C1-C6-alky1, ST, NR50C0R40, CONRSOR51, C1:G1)G2R41, etc.; R40 = H, OH, C1-C6-alky1, C2-C6-alkeny1, C2-C6-alkeny1, C1-C6-alky1, C2-C6-alkeny1, C2-C6-alkeny1, C1-C6-alky1, C1-C6-alky1, C2-C6-alkeny1, C2-C6-alkeny1, C1-C10-alky1, C1-C10-alkeny1, C2-C10-alkeny1, C2-C6-alkeny1, C2-C10-alkeny1, C1-C10-alkeny1, C1-C6-alky1;

R20 = haloalkyl, C2-C6-alkoxy, C2-C6-alkyl, alkylene-O-R8, alkylene-R8, alkylene(R8R9), alkylene = C1-C6-alkylene, R8, R9 = cycloalkyl, aryl, heterocyclyl, heteroaryl; Z = O, NR21, S, R21 = H, C1-C6-alkyl], were prepared to promote apoptosis. To illustrate the synthesis, reacting 3-02NC6H4OH with PhCH2CH2OH gave 3-02NC6H4OCH2CH2Ph which was reduced to the amine and acylated with 5-chlorosalicylic acid to give benzamide III.

821777-13-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of benzamide derivs, for promoting apoptosis)

RN 821777-13-3 ZCAPLUS

CN Benzenamine, N-(2-methoxyethyl)-N-methyl-4-nitro-2-(trifluoromethyl)- (CA INDEX NAME)

L70 ANSWER 3 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:802711 ZCAPLUS Full-text

DOCUMENT NUMBER: 141:314020

Preparation of substituted p-diaminobenzene TITLE:

derivatives as openers of the KCNQ family potassium

ion channels

INVENTOR(S): Khanzhin, Nikolay; Rottlaender, Mario; Ritzen,

Andreas; Watson, William Patrick

H. Lundbeck A/S, Den.

PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 176 pp.

CODEN: PIXXD2

Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

DOCUMENT TYPE:

PAT	ATENT NO.				KIND DATE				APPL	ICAT	ION I	NO.		D	ATE		
WO	2004				A1	-	2004	0930		WO 2	004-	DK18	6 6		2	0040	318 <
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		ES,	FΙ,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	SI,
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TD, TG																	
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CA 2519582		A1	A1 20040930				CA 2	004-	2519	582		2	0040	318 <			

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EP 1613303
                        A1 20060111
                                        EP 2004-721472
                                                               20040318 <--
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    CN 1761464
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                                         CN 2004-80007507
                                                               20040318 <--
                        Α
    JP 2006520759
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    IN 2005CN02347
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                                        IN 2005-CN2347
                                                               20050921 <--
                        Α
    NO 2005004848
                        Α
                            20051020
                                         NO 2005-4848
                                                               20051020 <---
    US 20060183791
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                              20060817
                                         US 2005-550448
                                                               20051116 <--
PRIORITY APPLN. INFO.:
                                         DK 2003-441
                                                            A 20030321 <--
                                         US 2003-456698P
                                                            P 20030321 <--
                                          WO 2004-DK186
                                                            W 20040318
OTHER SOURCE(S):
                      CASREACT 141:314020: MARPAT 141:314020
```

Y II X ZZLER3

- AB The title anilines I [s = 0-1; U = 0, S, SO2, etc.; q = 0-1; X = CO, SO2; with the proviso that q = 0 when X = SO2; Z = 0, S; R1 = H, alk(en/yn)yl, cycloalk(en)yl, etc.; R2 = H, alk(en/yn)yl, cycloalk(en)yl, etc.; R3 = alk(en/yn)yl, cycloalk(en)yl, etc.; Y = (un)substituted Ph, naphthyl, thienyl, etc.], useful for the prevention, treatment or inhibition of a disorder being responsive to an increased ion flow in a potassium channel, were prepared and formulated. Thus, reductive amination of Pr (4-amino-2-methylphenyl)carbamate (preparation given) with benzofuran-2-carbaidehyde in the presence of NaBH3CN afforded 43% Pr (4-(benzofuran-2-ylmethyl)amino-2-methylphenyl)carbamate. The compds. I have an ECO of <20000nM, in most cases <2000nM and in many cases <200nM in KCNQ2 channel assav.
- IT 335204-56-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of substituted p-diaminobenzene derivs. as openers of the KCNQ family potassium ion channels)

- RN 335204-56-3 ZCAPLUS
- CN Butanamide, 3,3-dimethyl-N-(2-methyl-4-nitrophenyl)- (CA INDEX NAME)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

2004:430796 ZCAPLUS Full-text

L70 ANSWER 4 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2004:430° DOCUMENT NUMBER: 141:7139

TITLE: Preparation of indolylquinoxalinones for treating hyperproliferative disorders and diseases associated

with angiogenesis

INVENTOR(S): Ladouceur, Gaetan H.; Bear, Brian; Bi, Cheng;

Brittelli, David R.; Burke, Michael J.; Chen, Gang; Cook, James; Dumas, Jacques; Sibley, Robert; Turner,

Michael R.

PATENT ASSIGNEE(S): Bayer Pharmaceuticals Corporation, USA

SOURCE: PCT Int. Appl., 217 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.												NO.			ATE			
		2004															0031	 110 <	<
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,	
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	GE,	
			GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	
			LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	
			OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	TJ,	TM,	
			TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	zw			
		RW:	BW,	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	
			BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	
			ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	
			TR,	BF,														TD,	
																	110 <		
		2003																	
	EP	1565	455			A1		2005	0824		EP 2	003-	7833	28		2	0031	110 <	<
		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,	
	BR	2003	0161	69		A		2005	0927		BR 2	003-	1616	9		2	0031	110 <	<
	CN	1738	814			A		2006	0222		CN 2	003-	8010	8639		2	0031	110 <	<
	MX	2005	0047	79		A													<
						A		2005	0609										
PRIO	RIT:	Y APP	LN.	INFO	.:														
																			<
																0030			
											WO 2	003-1	US36	003		W 2	0031	110	
	IE, SI, L BR 2003016169 CN 1738814 JP 2006509840 MX 2005004779 US 20060004011 NO 2005002796 RIORITY APPLN. INFO.:				.:	A T A A1 A		2005 2006 2006	0927 0222 0323 0722 0105 0609		BR 2 CN 2 JP 2 MX 2 US 2 US 2 US 2 US 2	003- 005- 005- 005- 005- 005- 002- 003-	1616 8010 5071 4779 5342 2796 4254 4609 4842	9 8639 46 15 90P 15P 02P		2 2 2 2 2 2 2 2 2 P 2 P 2	0031 0031 0031 0050 0050 0050 0050	110 < 110 < 504 < 506 609 < 112 < 407 <	< < <

OTHER SOURCE(S): MARPAT 141:7139

GI

$$\begin{bmatrix} R^1 & R^4 & R^3 & R^5 \\ Ax & R^2 & R^5 & R^5 \end{bmatrix}$$

AB The invention relates to title compds. I [wherein Ar = 6-membered aromatic ring containing 0-2 N atoms; R1 and R2 = independently H, halo, CF3, acyl, piperidinyl, piperazinyl, morpholinyl, or (un)substituted alkyl, alkoxy, amino, pyrrolidinyl, Ph. etc.; R3 = H. alkyl, OH, NO2, NH2, alkylamino, alkoxyamino, or (un) substituted benzoylamino; R4 = H, OH, halo, CN, acyl, sulfamovi, trialkylsiloxy, tetrazolyl, thienyl, pyrrolyl, pyrimidinyl, oxazolyl, furanyl, or (un)substituted alkyl, alkenyl, alkynyl, alkoxy, amino, oxadiazolyl, Ph, pyridyl(oxy), carbamoyl; R11 and R12 = independently H, F, or Cl with the proviso that when one of R11 and R12 = F or Cl, the other must be H; and pharmaceutically acceptable salts and esters thereof]. The invention also relates to the use of I and their pharmaceutical compns. for treating hyperproliferative disorders and diseases associated with angiogenesis (no data). Examples include representative syntheses for compds. of the invention, pharmaceutical compns, comprising them, and tumor model assays (no specific data given). For instance, N-Boc-indole was coupled with di-Me oxalate using t-BuLi to give tert-Bu 2-[methoxy(oxo)acetyl]-1H-indole-1carboxylate (72%). Cyclization of the dione with 1,2-phenylenediamine in AcOH afforded the quinoxalinone II (77%).

T 694533-16-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of indolylquinoxalinones for treating hyperproliferative disorders and diseases associated with angiogenesis) 694533-16-9 ZCAPLUS

RN 694533-16-9 ZCAPLUS
CN Benzenamine, N-(2-methoxyethyl)-N,3-dimethyl-4-nitro- (CA INDEX NAME)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 5 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2004:287838 ZCAPLUS Full-text

DOCUMENT NUMBER: 140:321373

TITLE: Preparation of novel pyrimidine amides as protein

kinase inhibitors

INVENTOR(S): Manley, Paul William; Breitenstein, Werner; Jacob,

Sandra; Furet, Pascal

PATENT ASSIGNEE(S): Novartis Ag, Switz.; Novartis Pharma GmbH SOURCE: PCT Int. Appl., 57 pp.

PCT Int. Appl., 57 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

E	PATENT NO.					KIN											ATE		
-		2004	0.20.0	20				2004				003					0020	026	,
V	40							AU,											
		w .						DK,											
								IN,											
								MN,											
								SY,											
				ZA,		50,	D.1.	017	10,	,	1117		,	0117	00,	00,	,,,	,	
		RW:				KG.	K7.	MD,	RII.	T.T.	TM.	AT.	BE.	BG.	CH.	CY.	CZ.	DE.	
								GB,											
			SI.	SK.	TR														
(CA	2499	822	,		A1		2004 2004 2007	0408		CA 2	003-	2499	822		2	0030	926	<
I	ΑU	2003	2702	77		A1		2004	0419		AU 2	003-	2702	77		2	0030	926	<
I	ΑU	2003	2702	77		B2		2007	0823										
		1546	127			A1		2005	0629		EP 2	003-	7506	39		2	0030	926	<
E	ΞP	1546	127			B1		2007	8080										
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
			ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK		
E	ЗR	2003	0147	97		A		2005	0726		BR 2	003-	1479	7		2	0030	926	<
(CN	1684	951			A		2005	1019		CN 2	003-	8232	13		2	0030	926	<
(CN	1004	0452	В		С		2005 2008 2006 2007	0723										
Ċ	JP	2006	5080	64		T		2006	0309		JP 2	004-	5390	39		2	0030	926	<
I	AΤ	3693	55			T		2007	0815		AT 2	003-	7506	39		2	0030	926	<
E	ΞS	2288	615			Т3		2008											
1	١Z	5389	30			A		2008											
2	ZA	5389: 2005: 2005: 2005: 8760:	0023	04		A		2006									0050		
1	·ίΧ	2005	0032	53		A		2005			MX 2	005-	3253			2	0050	323	<
	IN	2005	CN00	464		A		2007			IN 2	005-	CN46	4		2	0050	323	<
F	KR	8760.	55			B1		2008			KR 2	005-	7052	04		2	0050	325	<
		2005						2005									0050		
		1080						2008									0051		
		2006						2006			US 2	006-	5289	13		2	0060		
I	KR	2007	0989	40		A		2007	1005		KR 2	007-	7192	51		2	0070		
						A		2008	0125		IN 2	007-	CN43	30		- 2	0071	001	<
PRIOR	LTY	APP:	LN.	INFO	. :						GB 2	002-	2251	4		A 2	0020	927	<
																	0030		
											NK Z	005-	1052	04		AJ Z	0050	345	

OTHER SOURCE(S): MARPAT 140:321373

GI

$$\bigcap_{R \neq 1}^{R} \bigcap_{R \neq 1}^{R^2}$$

AB The title substituted N-(3-benzoylaminophenyl)-4-pyridyl-2-pyrimidinamines [I; R1 = H and R2 = NR5R6, or R1 = NR5R6 and R2 = H; R3 = alkyl, fluoroalkyl, hydroxyalkyl, carbamoyl; R4 = H, alkyl, halo; R5 and R6 = H, alkyl, hydroxyalkyl, etc. or NR5R6 = (un)substituted (un)saturated 5-7 membered ring optionally containing heteroatoms], useful for the therapy of a disease which responds to an inhibition of protein kinase activity, especially a neoplastic disease (e.g., leukemia), were prepared and formulated. Thus, amidation of 4-methyl-N-[4-(3-pyridinyl)-2-pyrimidinyl]-1,3-benzenediamine with 4-diethylamino-3-(trifluoromethyl)benzoic acid (preparation given) afforded I [R1 = H; R2 = NEt2; R3 = CF3; R4 = Me] which showed ICSO of 50-100 nM against c-Abl and ICSO of 200-500 nM against Bcr-Abl (in vitro inhibition data).

IT 677704-53-9P, 4-(Diethylamino)-3-(trifluoromethyl)benzonitrile
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)

(preparation of novel N-[3-(pyrimidin-2-ylamino)phenyl] benzamides as protein kinase inhibitors)

RN 677704-53-9 ZCAPLUS

CN Benzonitrile, 4-(diethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)

Ι



REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 6 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:591159 ZCAPLUS Fuil-text

DOCUMENT NUMBER: 139:157136

TITLE: Nile red type compound emitting red light, process for producing the same, and luminescent element utilizing

the same

INVENTOR(S): Nakaya, Tadao; Tajima, Akio; Saikawa, Tomoyuki;
Takano, Shinji; Yamauchi, Takao; Mori, Hidemasa

PATENT ASSIGNEE(S): Taiho Industries, Co. Ltd., Japan

SOURCE: PCT Int. Appl., 112 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 3 PATENT INFORMATION:

PATENT NO.		APPLICATION NO.	
WO 2003062213		WO 2003-JP477	
		BA, BB, BG, BR, BY,	
		DZ, EC, EE, ES, FI,	
GM, HR,	HU, ID, IL, IN, IS,	KE, KG, KP, KR, KZ,	LC, LK, LR, LS,
LT, LU,	LV, MA, MD, MG, MK,	MN, MW, MX, MZ, NO,	NZ, OM, PH, PL,
PT, RO,	RU, SC, SD, SE, SG,	SK, SL, TJ, TM, TN,	TR, TT, TZ, UA,
UG, US,	UZ, VC, VN, YU, ZA,	ZM, ZW	
RW: GH, GM,	KE, LS, MW, MZ, SD,	SL, SZ, TZ, UG, ZM,	ZW, AM, AZ, BY,
KG, KZ,	MD, RU, TJ, TM, AT,	BE, BG, CH, CY, CZ,	DE, DK, EE, ES,
FI, FR,	GB, GR, HU, IE, IT,	LU, MC, NL, PT, SE,	SI, SK, TR, BF,
BJ, CF,	CG, CI, CM, GA, GN,	GQ, GW, ML, MR, NE,	SN, TD, TG
JP 2003277371	A 20031002	JP 2002-14881	20020123 <
JP 2004018400	A 20040122	JP 2002-172127	20020612 <
JP 2003277369	A 20031002	JP 2003-12498	20030121 <
EP 1475372	A1 20041110	EP 2003-701142	20030121 <
R: AT, BE,	CH, DE, DK, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT,
		CY, AL, TR, BG, CZ,	
CN 1620441	A 20050525	CN 2003-802508	20030121 <
US 20050113575	A1 20050526	US 2004-501398	20040715 <
PRIORITY APPLN. INFO.	:	JP 2002-12222	A 20020121 <
		JP 2002-12224	A 20020121 <
		JP 2002-14881	A 20020123 <
		JP 2002-172127	
		JP 2001-313245	
		WO 2003-JP477	
OTHER SOURCE(S):	MARPAT 139:15713		20000422
GI		•	

R³ R⁴ N O O

- AB The invention relates to a Nile red-based red-emitting compound represented by I [Rl-2 = H and alkyl; R3(R5) = H and may combine with R1(R2) to form a ring; R4 = H and may combine with R3 to form a ring; X = H, halo, and -CH(CN)Ar]. The compound is suited for use as a red-emitting material in an organic light emitting device.
- IT 398482-49-0

RL: RCT (Reactant); RACT (Reactant or reagent)

(nile red type compound for red-emitting organic LED)

RN 398482-49-0 ZCAPLUS

CN Phenol, 5-[bis(1-methylethyl)amino]-2-nitro- (CA INDEX NAME)

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 7 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:511082 ZCAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 139:85343

TITLE: Preparation of 2-(heterocyclylmethyl)benzimidazoles as

respiratory syncytial virus antiviral agents

INVENTOR(S): Yu, Kuo-long; Wang, Xiangdong; Sun, Yaxiong; Cianci, Christopher; Thuring, Jan Willem; Combrink, Keith; Meanwell, Nicholas; Zhang, Yi; Civiello, Rita L.

PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: PCT Int. Appl., 149 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.			APPLICATION NO	
				0 20021206 <
WO 2003053344			110 2002 000322	0 20021200 (
			BA. BB. BG. BR. B	Y, BZ, CA, CH, CN,
				I, GB, GD, GE, GH,
			JP, KE, KG, KP, K	
				Z, NO, NZ, OM, PH,
				M, TN, TR, TT, TZ,
		VN, YU, ZA,		,,,
RW: GH, G	I. KE. LS. I	MW. MZ. SD.	SL. SZ. TZ. UG. Z	M, ZW, AM, AZ, BY,
				Z, DE, DK, EE, ES,
FI, F	, GB, GR,	IE, IT, LU,	MC, NL, PT, SE, S	I, SK, TR, BF, BJ,
CF, C	CI, CM,	GA, GN, GQ,	GW, ML, MR, NE, S	N, TD, TG
US 2003020786	A1	20031106	US 2002-309505	20021204 <
US 6919331	B2	20050719		
AU 2002362094	A1	20030709	AU 2002-362094	20021206 <
EP 1461035	A2	20040929	EP 2002-797226	20021206 <
R: AT, B	C, CH, DE,	DK, ES, FR,	GB, GR, IT, LI, L	U, NL, SE, MC, PT,
IE, S	, LT, LV, 1	FI, RO, MK,	CY, AL, TR, BG, C	Z, EE, SK
PRIORITY APPLN. IN	·o.:			P P 20011210 <
			WO 2002-US3922	0 W 20021206 <
OTHER SOURCE(S):	MARP	AT 139:85343	3	

60

AB Title compds. I [wherein R1 = (CRaRb)nX; R2 = H; R3 = CONRhRi, CO2Rd, or (un) substituted alkyl; R4 = NH2, CONRhRi, heteroaryl, alkenyl, CO2Rd, N=CPh2, C(NOH)NH2, C(NH)NH2, or (un)substituted alkyl; R5 = CO2Rj or (un)substituted alkyl or alkenyl; O = (un)substituted benzimidazolyl, benzotriazolyl, imidazopyridinyl, quinolinyl, quinazolinyl, benzyloxy, etc.; X = H or (un) substituted alkyl; Ra and Rb = independently H or (halo) alkyl; Rd = alkyl; Rh and Ri = independently H or alkyl; Rj = H or alkyl; n = 1-6; and pharmaceutically acceptable salts thereof] were prepared as antiviral compds. More particularly, the invention provides 2-(heterocyclylmethyl)benzimidazole derivs, for the treatment of respiratory syncytial virus (RSV) infection. For example, 1-isopropyl-1,3-dihydrobenzimidazol-2-one was coupled with 2chloromethyl-1-(3-methylbutyl)-1H-benzimidazole-5-carbonitrile in the presence of Cs2CO3 in DMF to give II (95%). Disclosed compds. protected HEp-2 cells from RSV-induced cytopathic effects with EC50 values between 50 μM and 0.001 μM , compared to an EC50 of 3 μM for ribavirin. I also displayed antiviral activity by reducing viral protein expression in HEp-2 cells with EC50 values between 50 μM and 0.001 μM , compared to an EC50 value of 3 μM for ribavirin. Thus, I and compns. comprising I are useful for the treatment of RSV infections.

IT 554457-74-8P 554458-11-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of (heterocyclylmethyl)benzimidazoles as RSV antiviral agents)

554457-74-8 ZCAPLUS

RN

CN Benzonitrile, 4-[(6-hydroxyhexyl)amino]-3-nitro- (CA INDEX NAME)

RN 554458-11-6 ZCAPLUS

CN Benzonitrile, 4-[(4-hydroxybutyl)amino]-3-nitro- (CA INDEX NAME)

L70 ANSWER 8 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2003:261620 ZCAPLUS Full-text

DOCUMENT NUMBER: 138:287673

TITLE: Preparation of phenylbenzimidazole compounds useful

for treating hepatitis C virus

INVENTOR(S): Priestley, Eldon Scott; Decicco, Carl P.; Hudyma,

Thomas W.; Zheng, Xiaofan

PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: PCT Int. Appl., 74 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.								APPL						ATE		
WO 200	30265	87		A2											0020	926 <-	
WO 200	30265	87		A3		2003	1106										
W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,	
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	
	PL,	PT,	RO,	RU,	SD,	SE.	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR.	TT.	TZ,	
	UA.	UG.	UZ.	VN.	YU.	ZA.	ZM.	ZW									
RV	GH,	GM.	KE.	LS.	MW.	MZ.	SD.	SL.	SZ.	TZ.	UG.	ZM.	ZW.	AM.	AZ.	BY.	
															EE,		
	FI, FR, G																
							GW,							,	,	,	
AU 200														2	0020	926 <-	
US 200																	
EP 142																	
	AT,																
Α.							MK,								PIC,	FI,	
US 200															0030	027 -	
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				BZ		2004	1012										
PRIORITY AF	PLN.	TMEO	• •													926 <-	
																926 <-	
											US30	989		W 2	0020	926 <-	
OTHER SOURC	E(S):			MARI	PAT	138:	2876	73									

62

AB Compds. of formula I [Q = CH, N; R1 = tetrazolyl, MeCONHSO2, PhCONHSO2, etc.; R2 = CH2-aryl, CHPh2, etc.; R3 = cycloalkyl] are prepared which are useful in treating viral hepatitis C. Thus, II was prepared and had an IC50 of 0.14 µM against HCV NS5B RdRp (RNA-dependent RNA polymerase).

IT 28096-55-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of phenylbenzimidazole compds. for treating hepatitis C viral infection)

RN 28096-55-1 ZCAPLUS

CN Benzonitrile, 4-(cyclohexylamino)-3-nitro- (CA INDEX NAME)

L70 ANSWER 9 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:418358 ZCAPLUS Full-text

DOCUMENT NUMBER: 137:370024

TITLE: Synthesis of some new

2-substituted-phenv1-1H-benzimidazole-5-carbonitriles

and their potent activity against Candida species
AUTHOR(S): Goker, Hakan; Kus, Canan; Boykin, David W.; Yildiz,

Sulhiye; Altanlar, Nurten

CORPORATE SOURCE: Faculty of Pharmacy, Department of Pharmaceutical

Chemistry, Ankara University, Ankara, 06100, Turk.

SOURCE: Bioorganic & Medicinal Chemistry (2002), 10(8),

2589-2596

CODEN: BMECEP: ISSN: 0968-0896

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 137:370024

AB New 2-substituted-phenyl-1H-benzimidazole-5-carboxylic acids, ethyl-5carboxylate, -5-carboxamides,-5-carboxaldehyde, -5-chloro-, -5-

trifluoromethyl, and -5-carbonitriles, -6-carbonitrile were prepared and evaluated in vitro against Candida species. The cyano substituted compds. exhibited the greatest activity with MIC values of 3.12 µg/mL, values similar to that of fluconazole.

T 28096-55-1P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (synthesis and m.p. of 2-substituted-Ph-1H-benzimidazole-5-carhonitriles)

RN 28096-55-1 ZCAPLUS

CN Benzonitrile, 4-(cyclohexylamino)-3-nitro- (CA INDEX NAME)



REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 10 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:122974 ZCAPLUS Full-text

DOCUMENT NUMBER: 136:191436

TITLE: Nile-red luminescent compound, process for producing the same, and luminescent element utilizing the same

INVENTOR(S): Nakaya, Tadao; Yamauchi, Takao; Saikawa, Tomoyuki; Tajima, Akio; Mori, Hidemasa

PATENT ASSIGNEE(S): Taiho Industries, Co., Ltd, Japan

KIND DATE

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1

DATENT INCODESTION.

PATENT INFORMATION:

PF	MIENI NO.	KIND	DATE AD	PEICATION NO.	DAIL
WC	2002012208	A1	20020214 WC	2001-JP5671	20010629 <
	W: CA, CN, JP, RW: AT, BE, CH, PT, SE, TR		, DK, ES, FI, E	FR, GB, GR, IE, IT	, LU, MC, NL,
		A1		2003-416586 2000-236670	20030513 < A 20000804 <
11/10//11	THE BIT. THE OT.		JE	2000-348516	A 20001115 <
OTHER S	SOURCE(S):	MARPAT	136:191436	2001-JP5671	W 20010629 <

ADDITORTION NO

DATE

AB The invention provides a novel nile-red compound capable of emitting a nearly crimson red light; a novel process for producing the compound; and a luminescent compound which emits a nearly crimson red color at a high luminance. The nile-red luminescent compound is obtained by converting C=O in a nile-red compound into C=CH-Ar (wherein Ar is a fluorinated aromatic group) and incorporating an electron-donating group into the nile-red skeleton. The novel production process comprises reacting a nile-red dve with an electronattracting aromatic MeCN. The nile-red based luminescent composition comprises a nile-red luminescent compound represented by I [R1 = R2 = C1-5 alkyl; R3 = H, or may be combined with R1 to form -CH2CH2-CR6R7- [R6 = R7 = H, and C1-5 alkyl, where the carbon in -CR6R7- is joined to the benzene ring]; R4 = H or may be combined with R3 to form naphthalene ring; R5 = H or may be combined with R2 to form -CH2CH2-CR8R9- [R8 = R9 = H and C1-5 alkvl, where the carbon in -CR8R9- is joined to the benzene ringl; Ar = fluorinated aroms.]. 398482-49-0

RL: RCT (Reactant); RACT (Reactant or reagent)

(production of nile-red luminescent compound for electroluminescent device) RN $\,$ 398482-49-0 $\,$ ZCAPLUS

CN Phenol, 5-[bis(1-methylethyl)amino]-2-nitro- (CA INDEX NAME)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 11 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2001:338479 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:353175

TITLE: Preparation of amides and ureas as activators of

soluble guanylate cyclase

INVENTOR(S): Selwood, David; Glen, Robert; Reynolds, Karen; Wishart, Grant

Wishart, Grant
PATENT ASSIGNEE(S): University College London, UK

SOURCE: PCT Int. Appl., 101 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT	ATENT NO.				KIND DATE			APPL	ICAT	ION I	NO.		D	ATE			
					-												
WO 2001	0326	04		A1		2001	0510		WO 2	000-	GB42	49		2	0001	106	<
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	
	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	
	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,	
	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	
	YU,	ZA,	ZW														
RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG 20010510 CA 2000-2389773 20001106 <--A1 EP 1237849 A1 20020911 EP 2000-973061 20001106 <--R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR 20030408 JP 2003513064 Т JP 2001-534758 20001106 <--PRIORITY APPLN. INFO .: GB 1999-26286 A 19991105 <--US 2000-201382P P 20000502 <--WO 2000-GB4249 W 20001106 <--

OTHER SOURCE(S): MARPAT 134:353175

AB The title compds. R4PZNR1R2 [I; R1, R2 = alkyl; R1R2 together form alkylene; Z = alkylene; P = a direct bond, X, Y, W, XY, YW, XYW (wherein W = O, S, NR3; R3 = H, alkvl; Y = UV; V = a direct bond, alkvlene; U = CS, CO, SO2, C(:NR); R = H, OH, alkyl; X = O, NR6; R6 = H, alkyl, alkenyl, etc.); R4 = alkyl, alkenyl, alkynyl, etc.], useful in the activation of soluble quanylate cyclase, were prepared E.g., synthesis of the urea II, starting with 4-bromoaniline and 1-(3-aminopropyl)pyrrolidine, was given. Biol. data for compds. I (e.g., IC50 for inhibition of platelet aggregation) were presented.

338981-22-9P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amides and ureas as activators of soluble quanylate cvclase)

338981-22-9 ZCAPLUS RN

CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 12 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2001:63979 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:100871

TITLE: Benzimidazolone derivatives, method of preparation and their use as phosphodiesterase inhibitors

INVENTOR(S): Sawada, Kozo; Inoue, Takayuki; Sawada, Yuki; Mizutani,

Tsuyoshi

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd., Japan SOURCE:

PCT Int. Appl., 159 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

								APPLICATION NO.										
								WO 2000-JP4687										
	W:						AU,											
							DM,											
							JP,											
							MN,											
				SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	
		ZA,																
	RW:						MZ,											
							GB,								SE,	BF,	ВJ,	
							GN,											
	2379																	
	AU 2000058531																	
EP	1196	391			A1		2002	0417		EP 2	000-	9444	21		2	0000	712	<
	R:						ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
							R0											
	2002																	
BR	BR 2000013041				A 20020716				BR 2000-13041					20000712 <				
	HU 2002002186													20000712 <				
	2002						2003											
	2003						2003											
								ZA 2002-29										
					A 20050311				IN 2002-KN19 MX 2002-340									
	2002															0020		
US	6582	351			B1		2003	0624								0020		
PRIORIT:	Y APP	LN.	INFO	.:						AU 1	999-	1747			A 1	9990	721	<
										AU 1								
										WO 2	000-	JP46	87		W 2	0000	712	<
OTHER SO	OURCE	(S):			MAR	PAT	134:	1008	71									

Benzimidazolone derivs. I, its prodrugs or pharmaceutically acceptable salts AB thereof, a method for their preparation, pharmaceutical compns. containing them, and usefulness in treatment or prevention of diseases mediated by cyclic quanosine-3',5'-monophosphate phosphodiesterase (cGMP-PDE) are claimed. In I, Xa = CH or N; ya = O, S; R1a = halogen, cyano, NO2 carbamoy1, lower alkylcarbamovl which may be substituted with a heterocyclic group, carboxy, protected carboxy, lower alkyl, halo(lower)alkyl, lower alkoxy, acyl, lower alkanesulfonyl. R2a = lower alkyl, cycloalkyl or heterocyclic group, among which the lower alkyl group may have 1-3 substituents = OH, protected OH, acvl, lower-alkoxy-substituted aralkyloxy, amino, lower alkylamino, acvlamino, lower alkoxycarbonylamino, lower alkanesulfonylamino, ureido, lower alkylureido, sulfamoylamino, protected carboxy, carboxy, lower alkanesulfonyl, lower alkylenedioxy, carbamoyl, lower alkyl carbamoyl and sulfamoyl; and the cycloalkyl group and the heterocyclic group may have 1-3 substituents = OH, protected OH, acvl, lower-alkoxy-substituted aralkyloxy, amino, acvlamino, lower alkoxycarbonylamino, lower alkanesulfonylamino, ureido, lower alkylureido, sulfamoylamino, protected carboxy, lower alkanesulfonyl, lower alkyl, hydroxy(lower)alkyl, protected hydroxy(lower)alkyl, lower alkylenedioxy, carbamoyl and sulfamoyl. R3a, R4a and R5a = same or different, H, halogen, lower alkanoyl, carboxy, protected carboxy, carbamoyl, nitro, cyano, lower alkyl optionally substituted by hydroxy, lower alkoxy or loweralkoxy-substituted aralkyl; or two of R3a, R4a and R5a may combine together to form a lower alkylenedioxy. M = 1, 2, provided that when R3a = H, R4a = lower alkoxy and R5a = H, halogen, cyano, lower alkyl, lower alkoxy, protected carboxy, carboxy or nitro, then (1) the lower alkyl for R2a has 1-3 substituents = OH, protected OH, acvl, lower-alkoxy-substituted aralkyloxy, amino, acylamino, lower alkoxycarbonylamino, lower alkanesulfonylamino, ureido, lower alkylureido, sulfamoylamino, protected carboxy, carboxy, lower alkanesulfonyl, lower alkylenedioxy, carbamoyl, lower alkyl carbamoyl and sulfamovl, (2) the cycloalkyl for R2a has 1-3 substituents = OH, protected OH, acyl, lower-alkoxy-substituted aralkyloxy, amino, acylamino, lower alkoxycarbonylamino, lower alkanesulfonylamino, ureido, lower alkylureido, sulfamovlamino, protected carboxy, lower alkanesulfonyl, lower alkyl, hydroxy(lower)alkyl, protected hydroxy(lower)alkyl, lower alkylenedioxy, carbamoyl and sulfamoyl, (3) the heterocyclic group for R2a = pyrrolidinyl, dioxanyl and piperidyl which groups may be substituted with protected carboxy, acvl, lower alkanesulfonyl, carbamovl or sulfamovl, (4) R1a = carbamovl, lower alkylcarbamoyl which may be substituted with a heterocyclic group, carboxy, protected carboxy, acvl. or lower alkanesulfonyl, (5) Xa = N; (6) m = 2; or

(7) yra = S. Pharmaceutical compns. containing the above compds. are claimed (with test data provided for 8 compds.) to be effective for treatment or prevention of diseases mediated by cGMP-PDE: angina, hypertension, pulmonary hypertension, congestive heart failure, glomerular diseases, renal tubulointestinal diseases, renal failure, atherosclerosis, conditions of reduced blood vessel patency, peripheral vascular disease, stroke, bronchitis, asthma, allergic rhinitis, urticaria, glaucoma, diseases characterized by disorders of gut motility, erectile dysfunction, female sexual dysfunction, impotence, diabetic complications, micturition disorder, or incontinence and storage of urine disorder. The method of preparation comprises reacting II with III (Z1 = halogen) in the presence of base. III are made by intramol. cyclization of IV (X = N). For example, to a solution of 1-(trans-4-hydroxycyclohexyl)-5trifluoromethyl-2,3-dihydro-1H- benzimidazol-2-one (200 mg) in anhydrous DMF (2 mL) was added portionwise NaH (29.3 mg, 60% dispersion in mineral oil) at 5° under N2 atmosphere, and the mixture was stirred at room temperature for 30 min. After adding 3,4-dimethoxybenzyl bromide (154 mg), the mixture was stirred at room temperature for 2 h. After workup, 3-(3,4-dimethoxybenzyl)-1-(trans-4- hydroxycyclohexyl)-5-trifluoromethyl-2,3-dihydro-1H-benzimidazol-2one (217.9 mg) was obtained as a colorless solid.

IIT 28096-55-19, 4-Cyclohexylamino-3-nitrobenzonitrile
320405-95-6P, 4-(1,1-Dimethyl-2-hydroxyethylamino)-3nitrobenzonitrile 320406-03-9P,
4-[((S)-1-Ethyl-2-hydroxyethyl)amino)-3-nitrobenzonitrile
320406-04-0P, 4-((S)-1-Ethyl-2-hydroxyethyl)aminol-3-

320406-04-0P, 4-[((R)-1-Ethyl-2-hydroxyethyl)amino]-3-nitrobenzonitrile

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; benzimidazolone derivs., method of preparation and use as phosphodiesterase inhibitors)

RN 28096-55-1 ZCAPLUS

CN Benzonitrile, 4-(cyclohexylamino)-3-nitro- (CA INDEX NAME)

RN 320405-95-6 ZCAPLUS

CN Benzonitrile, 4-[(2-hydroxy-1,1-dimethylethyl)amino]-3-nitro- (CA INDEX NAME)

RN 320406-03-9 ZCAPLUS

CN Benzonitrile, 4-[[(1S)-1-(hydroxymethyl)propyl]amino]-3-nitro- (CA INDEX NAME)

Absolute stereochemistry.

RN 320406-04-0 ZCAPLUS

CN Benzonitrile, 4-[[(1R)-1-(hydroxymethyl)propyl]amino]-3-nitro- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 13 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2000:401817 ZCAPLUS Fuil-text

13

DOCUMENT NUMBER: 133:30667

TITLE: Heteroaryl-containing thiourea derivatives useful as

inhibitors of herpes viruses
INVENTOR(S): Bloom, Jonathan David; Digrandi, Martin Joseph;

Dushin, Russell George; Lang, Stanley Albert; O'Hara,

Bryan Mark

PATENT ASSIGNEE(S): American Home Products Corporation, USA SOURCE: PCT Int. Appl., 164 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4 PATENT INFORMATION:

PA:	FENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	
					-												
WO	2000	0342	69		A1		2000	0615		WO 1	999-	US28:	892		19	9991	206 <
	W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,
		CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
		IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,
		SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	UZ,	VN,	YU,	ZA,	zw		
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
		DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,
		CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG				
US 6166028				A 20001226				US 1999-444782					19991122 <				
US	6197	803			B1		2001	0306		US 1	999-	4470	06		19	9991	122 <

US	6201013		B1	20010313	US	1999-4440	75	19991122	<
CA	2351390		A1	20000615	CA	1999-23513	390	19991206	<
EP	1140913		A1	20011010	EP	1999-96514	13	19991206	<
	R: AT,	BE, CH	H, DE,	DK, ES, FR,	GB, GI	R, IT, LI,	LU, NL, S	E, MC, PT	,
	IE,	SI, LT	LV,	FI, RO					
TR	200101598	3	T2	20011022	TR	2001-1598		19991206	<
BR	9916042		A	20011204	BR	1999-16042	2	19991206	<
HU	200100475	58	A2	20020429	HU	2001-4758		19991206	<
HU	200100475	58	A3	20020528					
JP	200253155	8	T	20020924	JP	2000-58671	16	19991206	<
AU	756043		B2	20030102	AU	2000-31122	2	19991206	<
US	6262082		B1	20010717	US	2000-66948	33	20000925	<
US	6271236		B1	20010807	US	2000-66994	13	20000926	<
ZA	20010043	73	A	20020918	ZA	2001-4373		20010528	<
NO	200100283	36	A	20010808	NO	2001-2836		20010608	<
MX	200100582	27	A	20010911	MX	2001-5827		20010608	<
BG 105580				20020131	BG	2001-10558	30	20010608	<
IN	2001KN008	365	A	20050311	IN	2001-KN865	5	20010822	<
IN	2001KN008	366	A	20050311	IN	2001-KN866	5	20010822	<
IN	2001KN008	368	A	20050311	IN	2001-KN868	3	20010823	<
US	200300366	553	A1	20030220	US	2002-99695	ō	20020315	<
US	6555561		B2	20030429					
PRIORITY	APPLN. I	INFO.:			US	1998-20854	10 A	19981209	<
					US	1998-15069	2P P	19981209	<
					US	1998-15069	98P P	19981209	<
					US	1998-15519	2P P	19981209	<
					US	1998-15524	10P P	19981209	<
					US	1998-20816	54 A	19981209	<
					US	1998-20855	59 A	19981209	<
					US	1998-20856	51 A	19981209	<
					US	1999-44478	32 A3	19991122	<
					WO	1999-US288	392 W	19991206	<
						2000-66953		20000926	<
						2001-KN492		20010508	
						2001-KN541		20010522	
					IN	2001-KN558	3 A3	20010525	<
OTHER CO	MIDOR (C).		M2 D1	DAT 122.2066	7				

OTHER SOURCE(S): MARPAT 133:30667

- AB Title compds. I and related compds. are disclosed [wherein R1-R5 = H, C1-6alkyl or perhaloalkyl, C2-6 alkenyl or alkynyl, C3-10 (hetero)cycloalkyl, (hetero)aryl, halo, CN, NO2, CO2R6, COR6, OR6, SR6, SOR6, SO2R6, CONR7R8, NR6N(R7R8), N(R7R8), or W-Y-(CH2)n-Z, provided that at least 1 of R1-R5 \neq H; or R2R3 or R3R4 form 3- to 7-membered heterocycloalkyl or heteroaryl fusion: R6, R7 = H, C1-6 alkyl or perhaloalkyl, or aryl; R8 = H, C1-6 alkyl or perhaloalkyl, C3-10 (hetero)cycloalkyl, (hetero)aryl; or NR7R8 forms 3- to 7membered heterocycloalkyl; A = heteroaryl; W = O, NR6, or bond; Y = CO, CO2, or bond; Z = C1-4 alkyl, CN, CO2R6, COR6, CONR7R8, OCOR6, NR6COR7, OCONR6, OR6, SR6, SOR6, SO2R6, SR6N(R7R8), N(R7R8) or Ph; G = arvl or heteroarvl; X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH)J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph, or PhCH2; n = 1-6]. I, or pharmaceutical salts thereof, are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), varicella-zoster virus (VZV), or (no data) Epstein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for approx. 25 compds. in 2-4 bioassays. For instance, the pyridinylthiazolecarboxamide derivative II had an IC50 of 0.001 ug/mL against HCMV wild-type in human foreskin fibroblast cell culture.
- IT 151951-35-8 338981-22-9 821776-85-6

821777-03-1 1098071-56-7

RL: PRPH (Prophetic)

(Heteroaryl-containing thiourea derivatives useful as inhibitors of herpes viruses)

- RN 151951-35-8 ZCAPLUS
- CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

- RN 338981-22-9 ZCAPLUS
- CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N2,N2-trimethyl- (CA INDEX NAME)

RN 821777-03-1 ZCAPLUS

CN 1,3-Propanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N3,N3-trimethyl- (CA INDEX NAME)

RN 1098071-56-7 ZCAPLUS

CN Benzenamine, 2-chloro-N-methyl-N-(2-methylpropyl)-4-nitro- (CA INDEX NAME)

$$\begin{array}{c} \text{C1} & \text{Me} \\ \text{N-Bu-i} \end{array}$$

IT 71145-96-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(starting material; preparation of heteroaryl-containing thiourea derivs.

inhibitors of herpes viruses)

RN 71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)



REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 14 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2000:401816 ZCAPLUS Full-text

DOCUMENT NUMBER: 133:30666

TITLE: Aryl- and heteroaryl-substituted thiourea derivatives

useful as inhibitors of herpes viruses

INVENTOR(S): Bloom, Jonathan David; Digrandi, Martin Joseph;

Dushin, Russell George; Lang, Stanley Albert; O'Hara, Bryan Mark

PATENT ASSIGNEE(S):

American Home Products Corporation, USA

SOURCE: PCT Int. Appl., 159 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	TENT :															ATE		
	2000						2000				999-					9991:	206	<
	W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,	
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		IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	
		SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	UZ,	VN,	YU,	ZA,	ZW			
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,	
		DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	
		CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG					
CA	2350	996			A1		2000	0615		CA 1	999-	2350	996		1	9991	206	<
BR	9915	993			A		2001	0904		BR 1	999-	1599	3		1	9991:	206	<
EP	1137	647			A1		2001	1004		EP 1	999-	9651	31		1	9991:	206	<
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
		ΙE,	SI,	LT,	LV,	FI,	RO											
HU	2001	0046	11		A2		2002	0429		HU 2	001-	4611			1	9991:	206	<
	2001																	
JP	2002	5315	57		T		2002	0924		JP 2	000-	5867	15		1	9991:	206	<
ZA	2001	0043	18		A		2002	0826		ZA 2	001-	4318			2	0010	525	<
NO	2001	0028	37		A		2001	0719		NO 2	001-	2837			2	0010	808	<
MX	2001	0058	35		A		2001	0911		MX 2	001-	5835			2	0010	808	<
PRIORITY	Y APP	LN.	INFO	. :						US 1	998-	2079	61		A 1	9981	209	<
										WO 1	999-	US28	838		W 1	9991	206	<
OTHER SO	DURCE	(S):			MAR	PAT	133:	30666	5									

- AB Title compds. I and related compds. and their pharmaceutical salts are disclosed [wherein A = heteroaryl; R1-R4 = H, C1-4 alkyl or perhaloalkyl, halo, C1-4 alkkoy, cyano; R1R2 or R3R4 = C5-7 aryl fusion; G = aryl or heteroaryl; and X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH)J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph or PhCR2; n = 1-6]. The compds. are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), and varicella-zoster virus (VZV), as well as (no data) Espetein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for approx. 35 compds. in 2-4 bioassays. For instance, the pyridine derivative II had an IC50 of 0.018 µg/mL against HCMV wild-type in human foreskin fibroblast cell culture.
- IT 151951-35-8 338981-22-9 821776-85-6 821777-03-1 1098071-56-7
 - RL: PRPH (Prophetic)
 - (Aryl- and heteroaryl-substituted thiourea derivatives useful as inhibitors of herpes viruses)
- RN 151951-35-8 ZCAPLUS
- CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

- RN 338981-22-9 ZCAPLUS
- CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N2,N2-trimethyl- (CA INDEX NAME)

RN 821777-03-1 ZCAPLUS

CN 1,3-Propanediamine, N1-(2-chloro-4-nitropheny1)-N1,N3,N3-trimethyl- (CA INDEX NAME)

RN 1098071-56-7 ZCAPLUS

CN Benzenamine, 2-chloro-N-methyl-N-(2-methylpropyl)-4-nitro- (CA INDEX NAME)

IT 71145-96-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(starting material; preparation of heteroaryl thiourea derivs. as inhibitors

of herpes viruses)

RN 71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 15 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2000:401809 ZCAPLUS Full-text

DOCUMENT NUMBER: 133:30657
TITLE: Heterocyc

Heterocyclic carboxamide-containing thiourea derivatives containing a substituted phenylenediamine group, useful as inhibitors of herpes viruses

INVENTOR(S): Bloom, Jonathan David; Curran, Kevin Joseph; Digrandi,

GI

Martin Joseph; Dushin, Russell George; Jones, Thomas Richard; Lang, Stanley Albert; Ross, Adma Antonia;

Terefenko, Eugene Anthony; O'Hara, Bryan Mark American Home Products Corporation, USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 159 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

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		2000											-US28				9991	206	<
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OTHER	SC	DURCE	(S):			MARI	PAT	133:	3065	7									

77

AB Title compds. I and related compds. are disclosed (wherein R1-R5 = H, C1-6 alkyl or perhaloalkyl, C2-6 alkenyl or alkynyl, C3-10 (hetero)cycloalkyl, (hetero)aryl, halo, CN, NO2, CO2R6, COR6, OR6, SR6, SOR6, SO2R6, CONR7R8, NR6N(R7R8), N(R7R8), or W-Y-(CH2)n-Z, provided that at least 1 of R1-R5 \neq H; or R2R3 or R3R4 form 3- to 7-membered heterocycloalkyl or heteroaryl fusion; R6, R7 = H, C1-6 alkyl or perhaloalkyl, or aryl; R8 = H, C1-6 alkyl or perhaloalkyl, C3-10 (hetero)cycloalkyl, (hetero)aryl; or NR7R8 forms 3- to 7membered heterocycloalkyl; R9-R12 = H, C1-4 alkyl or perhaloalkyl, halo, C1-4 alkoxy, or cyano, or R9R10 or R11R12 = C5-7 aryl fusion, provided that at least 1 of R9-R12 \neq H; W = O, NR6, or bond; Y = CO, CO2, or bond; Z = C1-4 alkyl, CN, CO2R6, COR6, CONR7R8, OCOR6, NR6COR7, OCONR6, OR6, SR6, SOR6, SO2R6, SR6N(R7R8), N(R7R8) or Ph; G = monocyclic heteroaryl; X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH)J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph, or PhCH2; n = 1-6]. I, or pharmaceutical salts thereof, are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), varicella-zoster virus (VZV), or (no data) Epstein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for 18 compds. in 4 bioassays. For instance, the N-(4-thioureidophenyl)furan-2-carboxamide derivative II had an IC50 of 0.4 µg/mL against HCMV wild-type in human foreskin fibroblast cell culture, and $0.5~\mu g/mL$ against HSV in an ELISA assay. ΙT 151951-35-8 338981-22-9 821776-85-6

821777-03-1 1098071-56-7

RL: PRPH (Prophetic)

(Heterocyclic carboxamide-containing thiourea derivatives containing a substituted phenylenediamine group, useful as inhibitors of herpes viruses)

RN 151951-35-8 ZCAPLUS

CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

$$\begin{array}{c|c} \text{C2N} & & \\ & \text{N-CH}_2\text{--}\text{CH}_2\text{--}\text{OMe} \\ \\ & \text{Me} \end{array}$$

- RN 338981-22-9 ZCAPLUS
- CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitropheny1)-N1,N2,N2-trimethy1- (CA INDEX NAME)

- RN 821777-03-1 ZCAPLUS
- CN 1,3-Propanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N3,N3-trimethyl- (CA INDEX NAME)

- RN 1098071-56-7 ZCAPLUS
- CN Benzenamine, 2-chloro-N-methyl-N-(2-methylpropyl)-4-nitro- (CA INDEX NAME)

- IT 71145-96-5
 - RL: RCT (Reactant); RACT (Reactant or reagent)
 (starting material; preparation of heterocyclic carboxamide-containing and
 phenylenediamine-containing thiourea derivs. as inhibitors of herpes

viruses)

RN 71145-96-5 ZCAPLUS
CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)

CN CF3

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 16 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2000:401808 ZCAPLUS Full-text

DOCUMENT NUMBER: 133:30588

TITLE: Alpha-methylbenzyl-containing thiourea derivatives

containing a phenylenediamine group, useful as

inhibitors of herpes viruses
INVENTOR(S): Bloom, Jonathan David; Curran, Kevin Joseph; Digrandi,

Martin Joseph; Dushin, Russell George; Lang, Stanley Albert; Norton, Emily Boucher; Ross, Adma Antonia;

O'Hara, Bryan Mark

PATENT ASSIGNEE(S): American Home Products Corporation, USA

SOURCE: PCT Int. Appl., 168 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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WO	2000	0342	60		A2		2000	0615										<
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PRIORITY	APPLN. INFO.:			US	1998-208902	A	19981209 <
				WO	1999-US28839	W	19991206 <

OTHER SOURCE(S): MARPAT 133:30588

AB Title compds. I and related compds. are disclosed [wherein R1-R5 = H, C1-6 alkyl or perhaloalkyl, C2-6 alkenyl or alkynyl, C3-10 (hetero)cycloalkyl, (hetero)arvl, halo, CN, NO2, CO2R6, COR6, OR6, SR6, SOR6, SO2R6, CONR7R8, NR6N(R7R8), N(R7R8), or W-Y-(CH2)n-Z, provided that at least 1 of R1-R5 \neq H; or R2R3 or R3R4 form 3- to 7-membered heterocycloalkyl or heteroaryl fusion; R6, R7 = H, C1-6 alkyl or perhaloalkyl, or aryl; R8 = H, C1-6 alkyl or perhaloalkyl, C3-10 (hetero)cycloalkyl, (hetero)aryl; or NR7R8 forms 3- to 7membered heterocycloalkyl; R9-R12 = H, C1-4 alkyl or perhaloalkyl, halo, C1-4 alkoxy, or cyano, or R9R10 or R11R12 = C5-7 aryl fusion; W = O, NR6, or bond; Y = CO, CO2, or bond; Z = C1-4 alkyl, CN, CO2R6, COR6, CONR7R8, OCOR6, NR6COR7, OCONR6, OR6, SR6, SOR6, SO2R6, SR6N(R7R8), N(R7R8) or Ph; G = aryl or fused bicyclic aryl; X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH)J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph, or PhCH2; n = 1-6]. I, or pharmaceutical salts thereof, are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), varicella-zoster virus (VZV), or (no data) Epstein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for approx. 320 compds. in 1-4 bioassays. For instance, the [[(phenylethyl)thioureido]phenyl]benzofurancarboxamide derivative II had an

[[(phenylethyl)thioureido]phenyl]benzofurancarboxamide derivative II had an IC50 of 1.3 µg/mL against HCMV wild-type in human foreskin fibroblast cell culture, and 0.10 µg/mL against VZV in an ELISA assav.

IT 71145-96-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(starting material; preparation of α -methylbenzyl-containing thiourea derivs, as inhibitors of herpes viruses)

RN 71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)



REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 17 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2000:401806 ZCAPLUS Full-text

DOCUMENT NUMBER: 133:30733

TITLE: Heterocyclic carboxamide-containing thiourea derivatives containing a phenylenediamine group,

useful as inhibitors of herpes viruses

INVENTOR(S): Bloom, Jonathan David; Curran, Kevin Joseph; Digrandi,

Martin Joseph; Dushin, Russell George; Jones, Thomas Richard; Lang, Stanley Albert; Ross, Adma Antonia; Terefenko, Eugene Anthony; O'Hara, Bryan Mark

PATENT ASSIGNEE(S): American Home Products Corporation, USA

SOURCE: PCT Int. Appl., 188 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4 PATENT INFORMATION:

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						ML,											
	6197																
	2350					0615											
	9916																
	1144							EP	19	99-	9630	23		1	9991	206	<
EP	1144																
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	2002																<
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	6271					0807											
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	2001			A		1001									0010		
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US	2003	0036	653			0220		US	20	02-	9969	5		2	0020	315	<
US	6555	561		B2	2003	0429											

PRIORITY APPLN. INFO.: US 1998-208559 A 19981209 <--US 1998-150692P P 19981209 <--US 1998-150698P 19981209 <--P US 1998-155192P P 19981209 <--US 1998-155240P P 19981209 <--US 1999-444782 A3 19991122 <--WO 1999-US28842 W 19991206 <--US 2000-669535 A3 20000926 <--

OTHER SOURCE(S): MARPAT 133:30733 GΙ

AB Title compds. I and related compds. are disclosed [wherein R1-R5 = H, C1-6 alkyl or perhaloalkyl, C2-6 alkenyl or alkynyl, C3-10 (hetero)cycloalkyl, (hetero)aryl, halo, CN, NO2, CO2R6, COR6, OR6, SR6, SOR6, SO2R6, CONR7R8, NR6N(R7R8), N(R7R8), or W-Y-(CH2)n-Z, provided that at least 1 of R1-R5 \neq H; or R2R3 or R3R4 form 3- to 7-membered heterocycloalkyl or heteroaryl fusion; R6, R7 = H, C1-6 alkyl or perhaloalkyl, or aryl; R8 = H, C1-6 alkyl or perhaloalkyl, C3-10 (hetero)cycloalkyl, (hetero)aryl; or NR7R8 forms 3- to 7membered heterocycloalkyl; W = O, NR6, or bond; Y = CO, CO2, or bond; Z = C1-4 alkyl, CN, CO2R6, COR6, CONR7R8, OCOR6, NR6COR7, OCONR6, OR6, SR6, SOR6, SO2R6, SR6N(R7R8), N(R7R8) or Ph; G = monocyclic heteroaryl; X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH)J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph, or PhCH2; n = 1-6]. I, or pharmaceutical salts thereof, are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), varicella-zoster virus (VZV), or (no data) Epstein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for approx. 350 compds. in 1-4 bioassays. For instance, the thioureidophenylthiadiazolecarboxamide derivative II had an IC50 of 0.0011

µg/mL against HCMV wild-type in human foreskin fibroblast cell culture.

151951-35-8 338981-22-9 821776-85-6 821777-03-1 1098071-56-7

RL: PRPH (Prophetic)

(Heterocyclic carboxamide-containing thiourea derivatives containing a phenylenediamine group, useful as inhibitors of herpes viruses)

151951-35-8 ZCAPLUS RN

CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

$$\begin{array}{c|c} \text{O2N} & & \\ & \text{N-} \text{ CH}_2\text{--} \text{ CH}_2\text{--} \text{ OMe} \\ \\ \text{C1} & & \text{Me} \end{array}$$

- RN 338981-22-9 ZCAPLUS
- CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitropheny1)-N1,N2,N2-trimethyl- (CA INDEX NAME)

- RN 821777-03-1 ZCAPLUS
- CN 1,3-Propanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N3,N3-trimethyl- (CA INDEX NAME)

- RN 1098071-56-7 ZCAPLUS
- CN Benzenamine, 2-chloro-N-methyl-N-(2-methylpropyl)-4-nitro- (CA INDEX NAME)

71145-96-5 ΙT

RL: RCT (Reactant); RACT (Reactant or reagent)

(starting material; preparation of heterocyclic carboxamide-containing thiourea

derivs. as inhibitors of herpes viruses)

71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)



REFERENCE COUNT:

1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 18 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN 2000:401786 ZCAPLUS Full-text

ACCESSION NUMBER:

DOCUMENT NUMBER: 133:30587

TITLE: Benzamide-containing aryl thiourea derivatives useful as inhibitors of herpes viruses

Bloom, Jonathan David; Curran, Kevin Joseph; Digrandi, INVENTOR(S): Martin Joseph; Dushin, Russell George; Lang, Stanley

Albert; Norton, Emily Boucher; Ross, Adma Antonia;

O'Hara, Bryan Mark

PATENT ASSIGNEE(S): American Home Products Corporation, USA

SOURCE: PCT Int. Appl., 169 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

PAT	TENT :	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE		
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WO	2000	0342.	38		A1		2000	0612		MO I	999-	J258	83/		1	999 I.	206 <-	-
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		IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	
		SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	UZ,	VN,	YU,	ZA,	ZW			
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,	
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BR 9916086 A 20010904 BR 1999-16086
EP 1137632 A1 20011004 EP 1999-963021
EP 1137632 B1 20040728
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TR 200101997

T2 20010029

HU 2001004763

A2 20020429

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T 20020924

JP 2000-86686

19991206 <---

AT 272032

T 20040815

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T 20041130

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US 1999-444782 A3 19991122 <--
US 1999-447006 A3 19991122 <--
US 1999-447006 A3 19991122 <--
US 1999-44706 A3 1999122 <--
        OTHER SOURCE(S): MARPAT 133:30587
```

86

AB Title compds. I and related compds. are disclosed (wherein R1-R5 = H, C1-6 alkyl or perhaloalkyl, C2-6 alkenyl or alkynyl, C3-10 (hetero)cycloalkyl, (hetero)arvl, halo, CN, NO2, CO2R6, COR6, OR6, SR6, SOR6, SO2R6, CONR7R8, NR6N(R7R8), N(R7R8), or W-Y-(CH2)n-Z, provided that at least 1 of R1-R5 \neq H; or R2R3 or R3R4 form 3- to 7-membered heterocycloalkyl or heteroaryl fusion; R6, R7 = H, C1-6 alkyl or perhaloalkyl, or aryl; R8 = H, C1-6 alkyl or perhaloalkyl, C3-10 (hetero)cycloalkyl, (hetero)aryl; or NR7R8 forms 3- to 7membered heterocycloalkyl; R9-R12 = H, C1-4 alkyl or perhaloalkyl, halo, C1-4 alkoxy, or cyano, or R9R10 or R11R12 = C5-7 aryl fusion, provided that at least 1 of R9-R12 \neq H; W = O, NR6, or bond; Y = CO, CO2, or bond; Z = C1-4 alkyl, CN, CO2R6, COR6, CONR7R8, OCOR6, NR6COR7, OCONR6, OR6, SR6, SOR6, SO2R6, SR6N(R7R8), N(R7R8) or Ph; G = aryl or fused bicyclic heteroaryl; X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH)J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph, or PhCH2; n = 1-6]. I, or pharmaceutical salts thereof, are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), varicella-zoster virus (VZV), or (no data) Epstein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for approx. 75 compds. in 2-4 bioassays. For instance, the thioureidophenylbenzamide derivative II had an IC50 of 1.5 µg/mL against HCMV wild-type in human foreskin fibroblast cell culture, and $0.04~\mu g/mL$ against HSV in an ELISA assav.

IT 151951-35-8 338981-22-9 821776-85-6 821777-03-1 1098071-56-7

RL: PRPH (Prophetic)

(Benzamide-containing aryl thiourea derivatives useful as inhibitors of herpes viruses)

RN 151951-35-8 ZCAPLUS

CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

- RN 338981-22-9 ZCAPLUS
- CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N2,N2-trimethyl- (CA INDEX NAME)

- RN 821777-03-1 ZCAPLUS
- CN 1,3-Propanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N3,N3-trimethyl- (CA INDEX NAME)

- RN 1098071-56-7 ZCAPLUS
- CN Benzenamine, 2-chloro-N-methyl-N-(2-methylpropyl)-4-nitro- (CA INDEX NAME)

- IT 71145-96-5
 - RL: RCT (Reactant); RACT (Reactant or reagent) (starting material; preparation of benzamide-containing aryl thiourea

derivs. as

inhibitors of herpes viruses)

RN 71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)

REFERENCE COUNT: 4

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 19 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2000:401785 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER:

133:30586

TITLE:

Acetamide and substituted acetamide-containing aryl thiourea derivatives useful as inhibitors of herpes viruses

INVENTOR(S):

Bloom, Jonathan David; Digrandi, Martin Joseph; Dushin, Russell George; Lang, Stanley Albert; O'Hara, Brvan Mark

PATENT ASSIGNEE(S):

American Home Products Corporation, USA

SOURCE: PCT Int. Appl., 159 pp.

CODEN: PIXXD2
Patent

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PA	TENT	NO.			KIN					APPL	ICAT	ION :	.00		D	ATE		
						-									-			
WC	2000	0342	37		A2		2000	0615		WO 1	999-	US28	844		1	9991	206	<
WC	2000	0342	37		A3		2000	1123										
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		CZ.	DE.	DK.	DM.	EE.	ES.	FI,	GB.	GD.	GE.	GH.	GM.	HR.	HU.	ID.	IL.	
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EF	1137																	
	R:	ΑT,						FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
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H	2001	0049	44		A2		2002	0429		HU 2	001-	4944			1	9991	206	<
H	2001	0049	44		A3		2003	0328										
JE	2002	5315	44		T		2002	0924		JP 2	000-	5866	85		1	9991	206	<
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MΣ	2001	0056	80		A		2001	0911		MX 2	001-	5680			2	0010	606	<
NO	2001	0028	34		А		2001	0807		NO 2	001-	2834			2	0010	608	<
PRIORIT				. :						US 1	998-	2083	16			9981		
											999-					9991		
										1	,,,	0020	0 1 1		1	J J J I	200	

- AB Title compds. I and related compds. are disclosed [wherein R1-R5 = H, C1-6 alkyl or perhaloalkyl, C2-6 alkenyl or alkynyl, C3-10 (hetero)cycloalkyl, (hetero)aryl, halo, CN, NO2, CO2R6, COR6, OR6, SR6, SOR6, SO2R6, CONR7R8, NR6N(R7R8), N(R7R8), or W-Y-(CH2)n-Z, provided that at least 1 of R1-R5 \neq H; or R2R3 or R3R4 form 3- to 7-membered heterocycloalkyl or heteroaryl fusion; R6, R7 = H, C1-6 alkyl or perhaloalkyl, or aryl; R8 = H, C1-6 alkyl or perhaloalkyl, C3-10 (hetero)cycloalkyl, (hetero)aryl; or NR7R8 forms 3- to 7membered heterocycloalkyl; R9-R12 = H, C1-4 alkyl or perhaloalkyl, halo, C1-4 alkoxy, or cyano, or R9R10 or R11R12 = C5-7 arvl fusion; W = O, NR6, or bond; Y = CO, CO2, or bond; Z = C1-4 alkyl, CN, CO2R6, COR6, CONR7R8, OCOR6, NR6COR7, OCONR6, OR6, SR6, SOR6, SO2R6, SR6N(R7R8), N(R7R8) or Ph; G = C1-6 alkyl; X = bond, NH, C1-6 alkyl, alkenyl, alkoxy, alkylthio, or alkylamino, or (CH) J; J = C1-6 alkyl, C3-7 cycloalkyl, Ph, or PhCH2; n = 1-6]. I, or pharmaceutical salts thereof, are useful in the treatment of diseases associated with herpes viruses, including human cytomegalovirus (HCMV), herpes simplex viruses (HSV), varicella-zoster virus (VZV), or (no data) Epstein-Barr virus, human herpesviruses-6 and -7, and Kaposi herpesvirus. Approx. 1000 example compds. prepared by standard methods are listed, with biol. data for approx. 160 compds. in 4 bioassays. For instance, the thioureidophenylacetamide derivative II had an IC50 of 0.8 ug/mL against HCMV wild-type in human foreskin fibroblast cell culture, and 2 ug/mL against HSV
- in an ELISA assay. IT 151951-35-8 338981-22-9 821776-85-6 821777-03-1 1098071-56-7
 - RL: PRPH (Prophetic)

(Acetamide and substituted acetamide-containing aryl thiourea derivatives useful as inhibitors of herpes viruses)

- RN 151951-35-8 ZCAPLUS
- CN Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX NAME)

- RN 338981-22-9 ZCAPLUS
- CN 1,3-Propanediamine, N3-(2-chloro-4-nitrophenyl)-N1,N1-dimethyl- (CA INDEX NAME)

- RN 821776-85-6 ZCAPLUS
- CN 1,2-Ethanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N2,N2-trimethyl- (CA INDEX NAME)

- RN 821777-03-1 ZCAPLUS
- CN 1,3-Propanediamine, N1-(2-chloro-4-nitrophenyl)-N1,N3,N3-trimethyl- (CA INDEX NAME)

- RN 1098071-56-7 ZCAPLUS
- CN Benzenamine, 2-chloro-N-methyl-N-(2-methylpropyl)-4-nitro- (CA INDEX NAME)

IT 71145-96-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(starting material; preparation of acetamide-containing aryl thiourea derivs. as

inhibitors of herpes viruses)

RN 71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)



REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 20 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2000:259962 ZCAPLUS Full-text

DOCUMENT NUMBER: 132:283910

TITLE: Agent and method for dyeing fibers

INVENTOR(S): Braun, Hans-Juergen; Semadeni, Pascal Andre

PATENT ASSIGNEE(S): Wella Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 56 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	TENT NO				KIN	D	DATE		P	PP	LI	CAT:	I NO	. OV		D	ATE		
	200002	2149	96		A1		2000		V	10	19	99-I	EP66	01		19	9990	908	<
	W: E RW: A		BE,		CY,	DE,	DK,	ES,	FI,	FR	,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	,
	198471 103759				A1 A1		2000 2000						1984 9446				9981		
EP	103759 R: A	AΤ,		CH,	B1 DE,	DK,	2004 ES,		GB,	GR	,	IT,	LI,	LU,	NL,	SE,	MC,	PT	
	990687	_			A		2000		_				5871						<
AT	200252 258782	2	59		T		2002 2004	0215	I	ΔT	19	99-9		24		19	990	908	
	221488 637940				T3 B1		2004 2002						9446: 58136	24 67			9990		<

PRIORITY APPLN. INFO.: DE 1998-19847192 A 19981013 <--WO 1999-EP6601 W 19990908 <--

OTHER SOURCE(S): MARPAT 132:283910

AB Direct dyes for fibers, especially hair, which contain 4-nitrophenylamine derivs. I [R1 = H, C1-4 alkyl; Y = II, cycloalkyl, NR2R3, (CH2)zOR4; ≥1 of R2, $R3 \neq H$; R4 = C1-4 alkyl; X = N, O; m = 1-3; n = 0, 1; z = 1-6] are prepared which are readily water soluble, fast towards light, friction, permanent waving, and perspiration, and produce a brilliant vellow coloration. I can be used in conjunction with other direct dyes or with oxidative dyes. Thus, 1fluoro-4-nitrobenzene reacted with 1-amino-2-methoxyethane to form N-(2methoxyethyl)-4-nitrophenylamine (III). A composition containing III 0.49, iso-PrOH 10.00, 28% aqueous lauryl alc. diglycol ether sulfate Na salt 10.00, and H2O to 100.00 g was adjusted to pH 5 or 8 and applied to bleached hair at 40° for 40 min to produce an intense lemon-vellow color at either pH.

263902-65-4P

RL: BUU (Biological use, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (agent and method for dyeing fibers)

RN 263902-65-4 ZCAPLUS

Benzenamine, N-(3-methoxypropy1)-2-methyl-4-nitro- (CA INDEX NAME) CM

MeO- (CH2)3-

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 21 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1998:125899 ZCAPLUS Full-text

DOCUMENT NUMBER: 128:250295

ORIGINAL REFERENCE NO.: 128:49441a,49444a

TITLE: Pattern recognition corresponding analysis of powder

frequency doubling effect of the nitrobenzene

derivatives

Yang, Rongsheng; Chen, Jiangzhong; Zhang, Hanhui AUTHOR(S): CORPORATE SOURCE:

Department of Chemistry, Fuzhou University, Fuzhou,

350002, Peop. Rep. China

SOURCE: Fuzhou Daxue Xuebao, Ziran Kexueban (1997), 25(3),

95-98

CODEN: FDXKEN; ISSN: 1000-2243

PUBLISHER . Fuzhou Daxue Xuebao Bianjibu

DOCUMENT TYPE: Journal

LANGUAGE: Chinese

AB The relationship between the structure and powder frequency doubling effect was studied by pattern recognition corresponding anal. method to avoid space group problems. Two groups of binding relationships were formulated, which can predict crystals with higher powder frequency doubling effect. Path for mol. engineering and crystal engineering was discussed.

204931-24-8

RL: PRP (Properties)

(pattern recognition corresponding anal. of powder frequency doubling effect of nitrobenzene derivative)

RN 204931-24-8 ZCAPLUS

CN 1-Propanol, 1-[[4-nitro-2-(trifluoromethyl)phenyl]amino]- (CA INDEX NAME)

L70 ANSWER 22 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1998:89624 ZCAPLUS Full-text

DOCUMENT NUMBER . 128 - 192539

ORIGINAL REFERENCE NO.: 128:38035a,38038a

TITLE: The reactivity of nitrophenyl-substituted cyclic

amines in dehydrogenations AUTHOR(S):

Moehrle, H.; Mehrens, J. CORPORATE SOURCE: Institut Pharmazeutische Chemie,

Heinrich-Heine-Universitaet, Duesseldorf, D-40225,

Germany

SOURCE: Zeitschrift fuer Naturforschung, B: Chemical Sciences

(1998), 53(1), 37-48

CODEN: ZNBSEN; ISSN: 0932-0776

PUBLISHER: Verlag der Zeitschrift fuer Naturforschung

DOCUMENT TYPE: Journal

LANGUAGE: German

CASREACT 128:192539 OTHER SOURCE(S):

1-(4-Nitrophenyl)-substituted piperidine and perhydroazepine were inert to Hα(II)-EDTA, while the corresponding α-pipecoline gave an amino ketone with ring cleavage. However, the corresponding 2-nitrophenyl compds. reacted to give a 2-piperidinone, an aminopentanal, and an aminohexanone, resp. By an addnl. substituent in 2'-position the 4-nitro compds. underwent dehydrogenation too. With a Me group resulted a pattern analogous to 2-nitro

products. A neighboring CH2OH function enhanced the reaction with formation of benzoxazines and if possible their further oxidized derivs., hydroxy lactams.

203510-10-5P

RL: SPN (Synthetic preparation); PREP (Preparation)

(dehydrogenation of nitrophenyl-substituted cyclic amines)

RN 203510-10-5 ZCAPLUS

Hexanal, 6-[(2-methyl-4-nitrophenyl)amino]- (CA INDEX NAME) CN

L70 ANSWER 23 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1996:561104 ZCAPLUS Fuil-text

DOCUMENT NUMBER: 125:275309 ORIGINAL REFERENCE NO.: 125:51485a,51488a

TITLE: Synthesis of substituted diphenylamines under phase

transfer catalysis
AUTHOR(S): Durantini, Edgardo N.; Chiacchiera, Stella M.; Silber,

Juana J.

CORPORATE SOURCE: Dep. Quim. Fis., Univ. Nacl. Rio Cuarto, Rio Cuarto,

5800, Argent.

SOURCE: Synthetic Communications (1996), 26(20), 3849-3858 CODEN: SYNCAV; ISSN: 0039-7911

PUBLISHER: Dekker
DOCUMENT TYPE: Journal

LANGUAGE: English
OTHER SOURCE(S): CASREAC

OTHER SOURCE(S): CASREACT 125:275309

GI

- AB A convenient procedure for the synthesis of N-[(trifluoromethyl)nitrophenyl] substituted anilines, e.g., I (Rl = H, R2 = NO2; Rl = NO2, R2 = H; R1 = H, R2 = cyano), by means of a chloro-substitution reaction under conditions of phase-transfer catalysis (PTC) is reported. The ipso-substitution product is obtained with high yield. This method provides a general procedure for the synthesis of diphenylamines bearing electron-withdrawing groups in both aromatic rinos.
- IT 175873-18-4P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of diphenylamines under phase transfer catalysis)

RN 175873-18-4 ZCAPLUS

CN Benzenamine, N-butyl-4-nitro-2-(trifluoromethyl)- (CA INDEX NAME)



L70 ANSWER 24 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1996:426976 ZCAPLUS Full-text

DOCUMENT NUMBER: 125:195846

125:36687a,36690a

ORIGINAL REFERENCE NO.: TITLE: Synthesis, some reactions and anti-ulcer activity of

some 2-amino-3-(substituted phenvl)selenazolidines Hornyak, Gyula; Feller, Antal; Lempert, Karoly AUTHOR(S): CORPORATE SOURCE: Res. Group Alkaloid Chem., Hungarian Academy Sci.,

Budapest, H-1521, Hung.

ACH - Models in Chemistry (1995), 132(6), 871-885 SOURCE:

CODEN: ACMCEI; ISSN: 1217-8969

PUBLISHER: Akademiai Kiado DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 125:195846 GI

- AB 2-Imino-3-(substituted phenyl)selenazolidine salts, e.g., I (R1 = H, NO2, CF3, X = Cl, Br), were prepared (1) by acid induced ring closure of N-(2selenocyanatoethyl)anilines, or (2) by fusion of anilines with 2bromoethylselenocyanate. E.g., 2-NO2C6H4NHCH2CH2SeCN is refluxed in dioxane the presence of ethanesulfonic acid to give I (R1 = H, HX = HO3SEt) in 93% yield. Diselenide, e.g., (ArNHCH2CH2Se)2, formation accompanying the syntheses according to Method 1 above was successfully suppressed. Some Nsubstituted derivs. (e.g., II, $R1 \neq R2 = C1$, NO2, Z = CHO, Ac, CONHPr, SO2Et) of selenazolidines I, as well as 3-aryl-selenazolidin-2-one III (X = Se), and its thiazolidinone analog III (X = S), were also prepared The gastroprotective (antiulcer) activity of some of I, II and III is reported. ΙT 180691-77-4P
 - RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and methylsulfonylation in the synthesis of amino(substituted phenyl)selenazolidines as antiulcer agents)

180691-77-4 ZCAPLUS

CN 1-Butanol, 2-[(2-chloro-4-nitrophenyl)amino]- (CA INDEX NAME)

L70 ANSWER 25 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1996:259463 ZCAPLUS Full-text

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

124:288977 124:53579a,53582a

TITLE:

Preparation of nitroarvlamines

INVENTOR(S): PATENT ASSIGNEE (S):

SOURCE:

Jautelat, Manfred Bayer A .- G., Germany Eur. Pat. Appl., 7 pp.

CODEN: EPXXDW DOCUMENT TYPE: Patent German LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PA:	TENT NO.			KINI) !	DATE	AP	PLICATI	ION NO.			DATE	
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	EP	695737			A1		19960207	EP	1995-1	111426			19950720	<
	EP	695737			B1		19970618							
		R: BE,	DE,	FR,	GB,	IT,	NL							
	DE	4427249			A1		19960208	DE	1994-4	1427249			19940802	<
	US	5684203			A		19971104	US	1995-5	504034			19950726	<
	CA	2154965			A1		19960203	ÇA	1995-2	2154965			19950728	<
	JP	08059571			A		19960305	JP	1995-2	211399			19950728	<
IC	RIT	APPLN.	INFO	. :				DE	1994-4	1427249	Z	4	19940802	<
HE	R S	DURCE (S):			CASI	REAC	T 124:288	3977; 1	MARPAT	124:2885	977			

OTHER SOURCE(S):

PRI

AB RNHZ(Xn)NO2 [R = H, (cyclo)alkyl, alkenyl, aryl, etc.; X = halo, cyano, alkyl, alkoxy, etc.: Z = (n+2)-valent (hetero)arvl group; n = 0-3) were prepared by amination of Z(Xn)NO2 [Z = (n+1)-valent (hetero)aryl group] by (RNH)2CY (Y = 0 or S) in the presence of a base, O, and a dipolar solvent. Thus, 3-(F3C)C6H4NO2 was heated 4h at 50° with (MeNH)2CO in DMSO containing NaOH and with air sparging to give 100% N-methyl-4-nitro-2-trifluoromethylaniline.

175873-18-4P, N-Butyl-4-nitro-2-trifluoromethylaniline RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(preparation of nitroarylamines)

175873-18-4 ZCAPLUS RN

Benzenamine, N-butvl-4-nitro-2-(trifluoromethvl)- (CA INDEX NAME)

L70 ANSWER 26 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1995:169712 ZCAPLUS Full-text

DOCUMENT NUMBER: 122:31106

ORIGINAL REFERENCE NO.: 122:6139a,6142a

TITLE: Preparation of p-phenylenediamines

INVENTOR(S): Urano, Fumyoshi; Kimura, Takahiro; Kametani, Miki

PATENT ASSIGNEE(S): Wako Pure Chem Ind Ltd, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF
DOCUMENT TYPE: Patent

LANGUAGE: Japanese

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 06219997	A	19940809	JP 1993-28593	19930125 <
PRIORITY APPLN. INFO.:			JP 1993-28593	19930125 <
OTHER SOURCE(S):	CASREA	CT 122:31106	; MARPAT 122:31106	
GI				

- AB The title compds. [I, R = NH2; R1 = C1-3 alkyl; R3 = H, C1-3 alkyl, C1-3 alkyn; X = O, NH; Z = H, COR4, SO2R4; R4 = C1-3 alkyl, (alkyl)phenyl; n = 1-3], useful as intermediates for azo dyes, pharmaceuticals, and agrochems. and photog. developers (no data), are prepared by nitration of I [R = H; Z = YR2; R2 = C1-3 alkyl, (alkyl)phenyl; Y = CO, SO2], treatment with H2O alkaline aqueous solns., and reduction of the resulting I [R = NO2; Z = H, COR4, SO2R4). I [R = H, R1 = Et, R3 = Me, XZ = OAc, n = 2) (preparation given) was nitrated by HNO3 and H2SO4 at -5 to 0° for 3 h and treated with aqueous NaOH in MeOH at 25-30° for 2 h to give 99% I [R = NO2, R1 = Et, R3 = Me, XZ = OH, n = 2), which was reduced by H with Pd/C in EtOH at 25-35° for 6 h to give 95% sulfate salt of I (R = NH2, R1 = Et, R3 = Me, XZ = OH, n = 2).
- RI: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of phenylenediamines from anilines via nitroanilines) RN 5217-13-6 ZCAPUS
- CN Ethanol, 2-[ethyl(3-methyl-4-nitrophenyl)amino]- (CA INDEX NAME)

L70 ANSWER 27 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1994:77020 ZCAPLUS Full-text 120:77020

DOCUMENT NUMBER:

ORIGINAL REFERENCE NO.: 120:13849a,13852a

Preparation of N-alkoxyalkyl p-phenylenediamine TITLE:

developers for oxidative hair dyes

INVENTOR(S): Rose, David; Lieske, Edgar; Hoeffkes, Horst

PATENT ASSIGNEE(S): Henkel K.-G.a.A., Germany

SOURCE: Ger. Offen., 7 pp.

CODEN: GWXXBX DOCUMENT TYPE: Patent

LANGUAGE: German FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

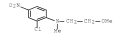
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 4205329	A1	19930826	DE 1992-4205329	19920221 <
WO 9316679	A1	19930902	WO 1993-EP351	19930213 <
W: JP, US				
RW: AT, BE, CH,	DE, DK	, ES, FR, GB	GR, IE, IT, LU, M	C, NL, PT, SE
PRIORITY APPLN. INFO.:			DE 1992-4205329	A 19920221 <
OTHER SOURCE(S):	MARPAT	120:77020		

GT

- AB Title compds. (I; R1, R2 = alkyli; X = H, halo; n = 2-4), were prepared Thus, 4-FC6H4NO2, CaCO3, and MeNHCH2CH2OMe were refluxed in acetone to give 4-O2NC6H4NMeCH2CH2OMe, which was hydrogenated followed by salification with HCl to give 4-H2NC6H4NMeCH2CH2OMe.2HCl. This developer together with 2-chloro-3amino-6methylphenol coupler and 3% H2O3 oxidizer gave a blue-black color to gray hair.
- ΤТ 151951-35-8P

RL: PREP (Preparation) (prepare of, as intermediate for developer for oxidative hair dves)

- RN 151951-35-8 ZCAPLUS
- Benzenamine, 2-chloro-N-(2-methoxyethyl)-N-methyl-4-nitro- (CA INDEX CN NAME)



L70 ANSWER 28 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1991:247193 ZCAPLUS Full-text

DOCUMENT NUMBER:

114:247193

ORIGINAL REFERENCE NO.: 114:41737a,41740a

TITLE:

Acyclic tertiary amines as nucleophiles in

substitution reactions of aromatic and heteroaromatic halides

AUTHOR(S):

Matsumoto, Kivoshi; Hashimoto, Shiro; Otani, Shinichi

CORPORATE SOURCE:

Coll. Lib. Arts Sci., Kyoto Univ., Kyoto, 606, Japan

Journal of the Chemical Society, Chemical SOURCE .

Communications (1991), (5), 306-7 CODEN: JCCCAT: ISSN: 0022-4936

Journal

DOCUMENT TYPE: LANGUAGE:

English



- AB Even acyclic tertiary amines such as Et3N, Pr3N and Bu3N, which have been believed to be inert to aromatic and heteroarom. halides, underwent SNAr reactions with aromatic and heteroarom, halides to give the dialkylamino derivs. via an addition-elimination mechanism; in the case of monocyclic amines like N-methylpyrrolidine (I) and N-methylpiperidine (II) the dealkylation being regioselective. Thus, the reaction of 2chlorobenzothiazole III (R = Cl) with I and II gave 92-96% III (R = pyrrolidino, piperidino) resp.
- 25900-35-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

- 25900-35-0 ZCAPLUS RN
- CN Benzenamine, N, N-diethyl-4-nitro-2-(trifluoromethyl)- (CA INDEX NAME)



L70 ANSWER 29 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1989:632859 ZCAPLUS Full-text DOCUMENT NUMBER: 111:232859

ORIGINAL REFERENCE NO.: 111:38689a,38692a

TITLE: Preparation and testing of

2,3(1H,4H)-quinoxalinediones as neuroleptics INVENTOR(S): Honore, Tage; Jacobsen, Poul; Elmelund, Flemming;

Naerum, Lars

PATENT ASSIGNEE(S): Aktieselskabet Ferrosan, Den.

SOURCE: Eur. Pat. Appl., 28 pp.

CODEN: EPXXDW DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.			APPLICATION NO.		DATE	
EP 315959	A2		EP 1988-118592			<
EP 315959						
EP 315959	B1	19940302				
R: AT, BE, CH	, DE, ES	, FR, GB, GE	R, IT, LI, LU, NL, SE			
DK 164317	c	19921102	DK 1987-5862		19871110	<
NO 8804729	A	19890511	NO 1988-4729		19881024	<
NO 179551	В	19960722				
NO 179551 NO 179551	С	19961030				
CA 1321587	C	19930824	CA 1988-582104		19881103	<
DK 8806206	A	19890511	DK 1988-6206		19881108	<
DK 8806206 DK 164317	В	19920609				
ZA 8808367	A	19890830	ZA 1988-8367		19881108	<
US 4948794		19900814				
AT 102192		19940315	AT 1988-118592		19881108	<
ES 2061606	Т3	19941216	ES 1988-118592		19881108	<
AU 8824949	A	19890511	AU 1988-24949		19881109	<
AU 618766	B2	19920109				
FI 8805151	A	19890511	FI 1988-5151		19881109	<
FI 100181	B1	19971015				
JP 01153680		19890615	JP 1988-282621		19881110	<
	B2	19980304				
KR 9711279	B1	19970709	KR 1988-14744		19881110	<
US 5026704	A	19910625	US 1989-456325		19891226	<
NO 9603412			NO 1996-3412		19960815	<
NO 307252	B1	20000306				
PRIORITY APPLN. INFO.:			DK 1987-5862	Α	19871110	<
				Α	19880316	<
			NO 1988-4729		19881024	
			EP 1988-118592		19881108	
			US 1988-268939	A1	19881108	<
OTHER SOURCE(S):	MARPAT	111:232859				
GI						

$$\begin{bmatrix} R^4 & R^5 & R^1 \\ R^3 & R^2 & R^3 \end{bmatrix}$$

- AB The title compds. [1] Rl = C3-8 cycloalkyl, aryl, aralkyl, (un)substituted C1-12 alkyl; R2-R5 = H, CF3 C1-4 alkoxy, cyano, halo, NO2; R2R3, R4R5 = atoms to complete an (un)substituted, fused aromatic ring; when Rl = Me, R3 ≠ CF3, MeO, Br, Cl, NO2] were prepared as neurotransmitter antagonists, especially at NMDA (N-methyl-D-aspartate), quisqualate, and kainate receptors. I are useful as tranquilizers. 2-Bromo-6-methoxynaphthalene was nitrated to give 73% 6-bromo-2-methoxy-1-nitronaphthalene which was treated with MeMH2 in DMF to give 95% 6-bromo-M-methyl-1-nitro-2-naphthylamine. The latter was hydrogenated over Pd-C and the product was refluxed with (CO2H)2.2H2O in 4 M HCl to give 92% benzoquinoxalinedione II. II inhibited AMPA (α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid) binding to quisqualate receptors in rat cerebral cortical membrane prepse, with an ICSO of 0.42 µg/mL.
- IT 28096-55-1P
 RI: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (preparation and reaction of, in preparation of tranquilizers)
 RN 28096-55-1 ZCAPPLUS
- CN Benzonitrile, 4-(cyclohexylamino)-3-nitro- (CA INDEX NAME)

L70 ANSWER 30 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1989:486773 ZCAPLUS Full-text

DOCUMENT NUMBER: 111:86773

ORIGINAL REFERENCE NO.: 111:14447a,14450a

TITLE: Organic crystals for nonlinear optics: molecular engineering of non-centrosymmetric crystal structures

AUTHOR(S): engineering of Nicoud, J. F.

CORPORATE SOURCE: Inst. Phys. Chim. Mater., Inst. Charles Sadron,

Strasbourg, F-67083, Fr.

SOURCE: Special Publication - Royal Society of Chemistry

(1989), 69 (Org. Mater. Non-linear Opt.), 157-62

CODEN: SROCDO; ISSN: 0260-6291

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Some recent results are presented concerning the use of chirality as an efficient strategy for the tentative control of non-centrosymmetry in organic crystals. In addition to the classical nitro-Ph or nitro-pyridine derivs., some new conjugated chains were studied, such as chalcone and 2-benzylidene-1,3-indanedione derivs. The nonlinear optical properties of these new materials were tested by 2nd-harmonic generation on crystalline powders, at

1.06 or 1.32 µm. II 115416-49-4 115416-50-7

RL: PROC (Process)
(second-harmonic generation of)

RN 115416-49-4 ZCAPLUS

CN 1-Propanol, 3-[[4-nitro-2-(trifluoromethyl)phenyl]amino]- (CA INDEX NAME)

RN 115416-50-7 ZCAPLUS

CN 1-Propanol, 3-[(4-nitro-3-(trifluoromethyl)phenyl]amino]- (CA INDEX NAME)

L70 ANSWER 31 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1988:463796 ZCAPLUS Full-text 109:63796

DOCUMENT NUMBER:

ORIGINAL REFERENCE NO.: 109:10551a,10554a

TITLE: Molecular and crystal engineering for organic

nonlinear optical materials AUTHOR(S): Nicoud, J. F.

CORPORATE SOURCE: ESPCI, UA, CNRS, Paris, 75231, Fr.

SOURCE: Molecular Crystals and Liquid Crystals (1988),

156(Pt. A), 257-68

CODEN: MCLCA5: ISSN: 0026-8941

DOCUMENT TYPE: Journal LANGUAGE: English

Nonlinear optical crystals are attractive due to their potential applications in optical signal processing. Several strategies were investigated to get efficient organic nonlinear optical crystals having enhanced quadratic responses and high efficiencies for 2nd harmonic generation. One chooses mols. with high mol. hyperpolarizability, β . Typical mols. possess a highly delocalized π-electron system bearing a donor and an acceptor group in such positions that an intramol. charge transfer occurs. These mols. must be assembled in an optimized acentric fashion to get a material having a high 2nd order elec. susceptibility, $\chi(2)$. The use of chirality to ensure noncentrosym. crystal structures, extra polar groups to influence dipoledipole interactions, push-pull N-oxide bonds and H bonding were satisfactory strategies to get some of the most efficient organic crystals for nonlinear optics.

115416-49-4P 115416-50-7P RL: PREP (Preparation)

(preparation of, in mol. and crystal engineering for organic nonlinear

optical materials)

RN 115416-49-4 ZCAPLUS

CN 1-Propanol, 3-[[4-nitro-2-(trifluoromethyl)phenyl]aminol- (CA INDEX NAME)

RN 115416-50-7 ZCAPLUS

CN 1-Propanol, 3-[[4-nitro-3-(trifluoromethyl)phenyl]amino]- (CA INDEX NAME)

L70 ANSWER 32 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1982:103817 ZCAPLUS Full-text

DOCUMENT NUMBER: 96:103817

ORIGINAL REFERENCE NO.: 96:17037a,17040a TITLE: Nitrating anilides

TITLE: Nitrating anilides
INVENTOR(S): Peer, Lydia; Mayer, Joseph

PATENT ASSIGNEE(S): Schering Corp., USA

SOURCE: U.S., 4 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4302599	A	19811124	US 1979-73838	19790910 <
PRIORITY APPLN. INFO.:			US 1979-73838	19790910 <

OTHER SOURCE(S): MARPAT 96:103817

Meta-substituted N-acylanilines reacted with HNO3-oleum mixts. at between -20 and +50° to yield p-nitro-N-acylanilines. Thus, 3-F3CC6H4NHCOCHMe2 was added slowly to oleum at .apprx.5°, 90% HNO3 was added dropwise, and the mixture was stirred 2 h at .apprx.5° to give 3,4-F3C(O2N)C6H3NHCOCHMe2. The products are useful as bactericides, herbicides, and antiandrogenic agents (no data). 80945-17-1P

IT 80

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 80945-17-1 ZCAPLUS

CN Pentanamide, 3-methyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L70 ANSWER 33 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1981:586828 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 95:186828

ORIGINAL REFERENCE NO.: 95:31169a,31172a

TITLE: N,N-Disubstituted p-phenylenediamine derivatives
INVENTOR(S): Wollemann, Bruno; Haertner, Harrtmut; Bardonner, Hans

PATENT ASSIGNEE(S): Merck Patent G.m.b.H. , Fed. Rep. Ger.

SOURCE: Ger. Offen., 17 pp.
CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2951100	A1	19810702	DE 1979-2951100	19791219 <
DE 2951100	C2	19830210		
FR 2471967	A1	19810626	FR 1980-26664	19801216 <
FR 2471967	B1	19840316		
GB 2065657	A	19810701	GB 1980-40329	19801217 <
GB 2065657	В	19830824		
JP 56095152	A	19810801	JP 1980-179116	19801219 <
JP 63053981	B	19881026		
US 4474987	A	19841002	US 1982-365131	19820405 <
PRIORITY APPLN. INFO.:			DE 1979-2951100 A	19791219 <
			US 1980-218249 A1	1 19801219 <
OTHER SOURCE(S):	MARPAT	95:186828		

AB Title phenylenediamines I (R = C1-6 alkyl; R1 = C1-6 alkyl, sulfo- or alkyleulfonamidoalkyl; R2 = H, alkyl), useful as developers for color photog. (no data), were prepared Thus, 3-MeC6H4NEtCH2CH2OH was nitrosated with Me2CHONO, and the nitroso compound hydrogenated over Pd/C to give II (as the sulfate monohydrate).

ΤТ 52177-13-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and hydrogenation of)

52177-13-6 ZCAPLUS RN

CN Ethanol, 2-[ethyl(3-methyl-4-nitrophenyl)amino]- (CA INDEX NAME)

L70 ANSWER 34 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1981:575793 ZCAPLUS Full-text

DOCUMENT NUMBER: 95:175793

ORIGINAL REFERENCE NO.: 95:29269a,29272a

TITLE: Aniline derivatives as anti-testosterone drugs

PATENT ASSIGNEE(S): Scherico Ltd., Switz.

SOURCE: Jpn. Kokai Tokkyo Koho, 18 pp.

CODEN: JKXXAF DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 56063916	A	19810530	JP 1980-87231	19800626 <
JP 56049884	В	19811125	JP 1971-45301	19710624 <
JP 58140052	A	19830819	JP 1982-161540	19820916 <
PRIORITY APPLN. INFO.:			JP 1971-45301 A	19710624 <
			DE 1971-2130450 A	19710619 <
			JP 1980-87231	19800626 <

P GI

I (R = H or C1-4 alkvl, X = NO2 or C1, Y = H, Me, CF3, F, or C1, Z = H, AB COCHMe2, or other acyl groups) are synthesized as testosterone inhibitors. Thus, N-(isobutyryl)-4-nitro-3-trifluoromethylaniline (II) [13311-84-7] was prepared by treating 4-nitro-3-trifluoromethylaniline [393-11-3] with isobutyryl chloride [79-30-1]. Tablets were prepared by combining II 5, starch 5, lactose 89.5, and Mg stearate 0.5 mg/tablet. 39240-77-2P TT

RL: PREP (Preparation)
(preparation of, as testosterone inhibitor)

RN 39240-77-2 ZCAPLUS

CN Butanamide, 2,3-dimethyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

L70 ANSWER 35 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1980:58461 ZCAPLUS Full-text

DOCUMENT NUMBER: 92:58461

ORIGINAL REFERENCE NO.: 92:9675a,9678a
TITLE: Cvcloalkanecar

TITLE: Cycloalkanecarboxanilide derivative herbicides
INVENTOR(S): Pilgram, Kurt H. G.; Skiles, Richard D.

PATENT ASSIGNEE(S): Shell Oil Co., USA SOURCE: U.S., 5 pp.

SOURCE: U.S., 5 pp.
CODEN: USXXAM
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4168153	A	19790918	US 1978-876594	19780210 <
CA 1087186	A1	19801007	CA 1978-294281	19780104 <
BE 863074	A1	19780719	BE 1978-184445	19780119 <
SE 7800692	A	19780722	SE 1978-692	19780119 <
NL 7800656	A	19780725	NL 1978-656	19780119 <
DE 2802282	A1	19780727	DE 1978-2802282	19780119 <
JP 53092739	A	19780815	JP 1978-3778	19780119 <
BR 7800354	A	19781010	BR 1978-354	19780119 <
AU 7832543	A	19790726	AU 1978-32543	19780119 <
AU 523765	B2	19820812		
AT 7800395	A	19800615	AT 1978-395	19780119 <
AT 360799	В	19810126		
GB 1593932	A	19810722	GB 1978-2214	19780119 <
CH 637917	A5	19830831	CH 1978-563	19780119 <
PRIORITY APPLN. INFO.:			US 1977-761515	A2 19770121 <
CT.				

RR1N NHCO R2

- AB Seventeen cyclopropanecarboxanilides I (R = H, Me, Pr; R1 = H, Me, Me2CH, Me3C, cyclopropyl, l-methylcyclopropyl, Pr; R2 = Me, Cl, Et; X = CF3, NO2, Br, Me) were prepared by acylation of the appropriate aniline with a cyclopropanecarbonyl chloride. Tests with 15 I showed both pre- and postemergence herbicidal activity. Thus, acylation of 4,3- (Me2CHNH) (F3C)C6H3NH2 with 1-methylcyclopropanecarbonyl chloride gave 84% I (R = H, R1 = Me2CH, R2 = Me, X = CF3).
- IT 70339-08-1P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation and reduction of nitro group in)
- RN 70339-08-1 ZCAPLUS
 CN Benzenamine, N-(1-methylethyl)-4-nitro-2-(trifluoromethyl)- (CA INDEX NAMF)



L70 ANSWER 36 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1979:523221 ZCAPLUS Full-text

DOCUMENT NUMBER: 91:123221

ORIGINAL REFERENCE NO.: 91:19879a,19882a

TITLE: Dipole moment study of the ortho-effect in

4-substituted NN-dimethyl-2-trifluoromethylanilines AUTHOR(S): Hallas, Geoffrey; Hepworth, John D.; Jones, Peter;

Ibbitson, Douglas A.; Jones, Alan M.; Turton, Andrew R.

CORPORATE SOURCE: Dep. Colour Chem., Leeds Univ., Leeds, UK

SOURCE: Journal of the Chemical Society, Perkin Transactions 2: Physical Organic Chemistry (1972-1999) (1979),

(4), 525-8

CODEN: JCPKBH; ISSN: 0300-9580

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Apparent dipole moments of 4,2-R(CF3)G6H3NMe2 (R = H, Me, Cl, Br, iodo, cyano, O2N, Me2N, H2N) in C6H6 were determined Vector moments in the direction of the major axis of the ring were calculated to explain the observed dipole moments. These vector moments are compared with corresponding values calculated for 4-RC6H4NMe2 (R = Cl, Br, iodo, cyano, O2N). The influence of the 2-CF3 group on the extent to which para substitution restores conjugation between the Me2N group and the ring is noted.

IT 71145-96-5P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation and ortho effect in, dipole moment study of) RN 71145-96-5 ZCAPLUS

MM /1143-96-3 ZCAPEOS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)



L70 ANSWER 37 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1979:490927 ZCAPLUS Full-text

DOCUMENT NUMBER: 91:90927

ORIGINAL REFERENCE NO.: 91:14683a,14686a

TITLE: Evidence for electronic buttressing in 4-substituted

NN-dimethyl-2-trifluoromethylanilines: long range

fluorine-19-proton coupling

AUTHOR(S): Bartle, Keith D.; Hallas, Geoffrey; Hepworth, John D.;

Jones, Peter; Matthews, Raymond S.

CORPORATE SOURCE: Dep. Phys. Chem., Univ. Leeds, Leeds, UK

SOURCE: Journal of the Chemical Society, Perkin Transactions
2: Physical Organic Chemistry (1972-1999) (1979),

2: Physical Organic Chemistry (1972-1999) (1979), (2), 192-4

CODEN: JCPKBH; ISSN: 0300-9580

DOCUMENT TYPE: Journal

LANGUAGE: English

AB The 1H NMR spectra of fourteen 4,2-R(F3C)C6H4NMe2 were examined A long range 19F-IH coupling 5J(F,H) between NMe and CF3 was confirmed by 19F NMR. A linear relation holds between 5J(F,H) in these compds. and the appropriate Hammett para substituent consts. in accord with the operation of electronic buttressing; a similar correlation holds between the chemical shift of the N-Me protons and op.

IT 71145-96-5

RL: PRP (Properties)

(NMR of, electronic buttressing in relation to)

RN 71145-96-5 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-3-(trifluoromethyl)- (CA INDEX NAME)



L70 ANSWER 38 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1979:203718 ZCAPLUS Full-text

DOCUMENT NUMBER: 90:203718

ORIGINAL REFERENCE NO.: 90:32397a,32400a

TITLE: Cycloalkanecarboxanilides with herbicide activity

INVENTOR(S): Pilgram, Kurt H. G.; Skiles, Richard D.

PATENT ASSIGNEE(S): Shell Internationale Research Maatschappij B. V.,

Neth. SOURCE: Fr. D

Fr. Demande, 57 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Patent
LANGUAGE: French
FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE	
FR 2377999	Al	19780818	FR 1978-1493		19780119	<
FR 2377999	B1	19810327				
CA 1087186	A1	19801007	CA 1978-294281		19780104	<
BE 863074	A1	19780719	BE 1978-184445		19780119	<
SE 7800692	A	19780722	SE 1978~692		19780119	<
NL 7800656	A	19780725	NL 1978-656		19780119	<
DE 2802282	A1	19780727	DE 1978-2802282		19780119	<
JP 53092739	A	19780815	JP 1978-3778		19780119	<
BR 7800354	A	19781010	BR 1978-354		19780119	<
AU 7832543	A	19790726	AU 1978-32543		19780119	<
AU 523765	B2	19820812				
AT 7800395	Ä	19800615	AT 1978-395		19780119	<
AT 360799	В	19810126				
GB 1593932	A	19810722	GB 1978-2214		19780119	<
CH 637917	A5	19830831	CH 1978~563		19780119	<
RITY APPLN. INFO.:			US 1977-761515	A	19770121	<

AB Minety-four herbicidal I (R = C1-6 alkyl, C1, F, Br, alkoxy or alkylthic; R1 = halo, CN, NO2, CR3:NOR3, COR3, Q1R3; R2 = optionally halogenated alkyl, alkenyl or aryl, alkynyl, alkoxyalkyl, etc.; R3 = H, alkyl or cycloalkyl; Q = O, S, SO, SO2 or NR3; O1 = O, S, SO or SO2) were prepared; extensive data for both pre- and postemergence application are given. Thus, 3,4-(F3C)C1C6H3NO2 was treated with Me2CHONa, the NO2 group was reduced, and the aniline was acylated with 1-methylcyclopropanecarbonyl chloride to give I (R = Me, R1 = C1, R2 = Me2CH, Q = O).

IT 70339-08-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reduction of)

RN 70339-08-1 ZCAPLUS

CN Benzenamine, N-(1-methylethyl)-4-nitro-2-(trifluoromethyl)- (CA INDEX NAME)

L70 ANSWER 39 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1978:22155 ZCAPLUS Full-text
DOCUMENT NUMBER: 88:22155

DOCUMENT NUMBER: 88:22155 ORIGINAL REFERENCE NO.: 88:3549a,3552a

TITLE: Alkanoic acid anilides

INVENTOR(S): Neri, Rudolph O.; Topliss, John G.

PATENT ASSIGNEE(S): Scherico Ltd., Switz.

SOURCE: Can., 56 pp.
CODEN: CAXXA4

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CA 1012462	A1	19770621	CA 1971-116108	19710618 <
PRIORITY APPLN. INFO.:			CA 1971-116108 A	19710618 <
GT				

$$R_2$$
CHCHMeC(Z)NR1 R_2

- AB Eight alkanoic anilides I (Z=0, S; R=H, Me; R1=H, Me, EI; R2=CF3, C1, Br; R3=NO2, C1), which are useful as anti-androgenic agents (no data), were prepared by various methods. Amidation of Me2CHCOCl by an aniline derivative yielded I (Z=0, R=R1=H, R2=CF3, R3=NO2). The Beckmann rearrangement of 4,3-Cl(R3CC)CGH3C(SCH3C(NOH)CHMe2 gave I (R=R1=H, R2=CF3, R3=C1, Z=O). I (R=R1=H, R2=CF3, R3=NO2, Z=S) was prepared from Me2CHCS2Me and the resp. aniline.
- IT 39240-77-2P
 - RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 39240-77-2 ZCAPLUS
- CN Butanamide, 2,3-dimethyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

L70 ANSWER 40 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1977:189358 ZCAPLUS Full-text

DOCUMENT NUMBER: 86:189358

ORIGINAL REFERENCE NO.: 86:29689a,29692a

TITLE: Synthesis of 4-aminodiphenylamine and its relatives

AUTHOR(S): Rondestvedt, Christian S., Jr.

CORPORATE SOURCE: Jackson Lab., E. I. Du Pont de Nemours and Co.,

Wilmington, DE, USA

SOURCE: Journal of Organic Chemistry (1977), 42(10), 1786-90

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 86:189358

CT

AB Formanilides and acetanilides condensed with halonitrobenzenes I (R = NO2, H, Cl; R2 = H, CF3; R3 = H, Me, NO2; X = Cl, F) to give eight resp. diohenvlamines II (R3 = H, Me: R4 = H, OBt).

IT 61587-18-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 61587-18-6 ZCAPLUS

CN Benzenamine, N-cvclohexvl-4-nitro-3-(trifluoromethvl)- (CA INDEX NAME)

L70 ANSWER 41 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1975:409343 ZCAPLUS Full-text

DOCUMENT NUMBER: 83:9343

ORIGINAL REFERENCE NO.: 83:1545a,1548a

TITLE: Substituted N,N-dimethylanilines containing an o-trifluoromethyl group. Synthesis of

N.N-dimethyl-2-trifluoromethylaniline

AUTHOR(S): Hepworth, John D.; Jones, Peter; Hallas, Geoffrey

CORPORATE SOURCE: Div. Chem., Polytech, Preston, Preston, UK

SOURCE: Synthesis (1974), (12), 874-6 CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE: Journal LANGUAGE: English

LANGUAGE: English
OTHER SOURCE(S): CASREACT 83:9343

GI For diagram(s), see printed CA Issue.

AB The amines I (X = Me2N, MeNH, piperidino, morpholino, Et2N) were prepared in 74-94% yield by the reaction of I (X = Cl) with the corresponding amine. Reduction of I (X = Me2N) gave 94% 4,2-H2N(F3C)C6H3NMe2 (II), which was converted to its Ac and Bz derivs. Diazotization of II followed by treatment with CuBr in HBr gave 91% 4,2-Br (F3C)C6H3NMe2, which when treated with BuLi gave 72% 2-F3CC6H4NMe2.

IT 25900-35-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 25900-35-0 ZCAPLUS

CN Benzenamine, N.N-diethvl-4-nitro-2-(trifluoromethvl)- (CA INDEX NAME)



L70 ANSWER 42 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1974:425336 ZCAPLUS Full-text 81:25336

DOCUMENT NUMBER:

ORIGINAL REFERENCE NO.: 81:4085a,4088a

TITLE: p-Nitroaniline derivatives INVENTOR(S):

Suda, Hideaki; Kanda, Tatsuo; Tomita, Hiroshige; Nakanishi, Hirotoshi; Hida, Hiroshi; Nuno, Tatsumi

PATENT ASSIGNEE(S): Sumitomo Chemical Co., Ltd.

Jpn. Kokai Tokkyo Koho, 4 pp. SOURCE:

CODEN: JKXXAF DOCUMENT TYPE: Patent

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
JP 49020129	A	19740222	JP 1972-63507		19720623 <
CA 1012553	A1	19770621	CA 1973-174587		19730621 <
DE 2331900	A1	19740124	DE 1973-2331900		19730622 <
DE 2331900	C2	19861113			
FR 2199533	A1	19740412	FR 1973-22956		19730622 <
IT 991637	В	19750830	IT 1973-68879		19730622 <
GB 1421600	A	19760121	GB 1973-29902		19730622 <
CH 593236	A5	19771130	CH 1973-9211		19730625 <
FR 2225423	A1	19741108	FR 1974-3932		19740206 <
FR 2225423	B1	19780616			
PRIORITY APPLN. INFO.:			JP 1972-18191	A	19720213 <
			JP 1972-63507	A	19720623 <
			JP 1972-63508	A	19720623 <
			JP 1972-63509	A	19720623 <
			JP 1972-63512	A	19720623 <
			JP 1972-63513	Α	19720623 <
			JP 1972-64714	Α	19720627 <
			JP 1972-66286	A	19720701 <
			JP 1973-18191	Α	19730213 <

GI For diagram(s), see printed CA Issue.

AB The nitroanilines (I; R1, R2 = H, C1-6alkyl, CH2CH2OH, CH2CH2Cl, β acvlaminoethyl, CH2CH2NHSO2Me, but not R1 = R2 = H) were prepared by treating the nitrophenol ethers (II; R3 = C1-6 alkyl, Ph) with HNR1R2. Thus, heating II (R3 = Me) with 15 mole 20% agueous EtNH2 at 180° for 10 hr gave 97% I (R1 =Et, R2 = H) with 93% conversion. Among 5 more I prepared were the following (R1, R2 given): Pr,H; CH2CH2OH,H; CH2CH2OH,Et; CH2CH2NHSO2Me, Et.

52177-13-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 52177-13-6 ZCAPLUS

CN Ethanol, 2-[ethyl(3-methyl-4-nitrophenyl)aminol- (CA INDEX NAME)

L70 ANSWER 43 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1974:108165 ZCAPLUS Full-text

DOCUMENT NUMBER: 80:108165

ORIGINAL REFERENCE NO.: 80:17387a,17390a

TITLE:

N, N-Disubstituted p-phenylenediamines

INVENTOR(S): Suda, Hideaki; Kanda, Tatsuo; Tomita, Hiroshige; Nakanishi, Hirotoshi; Hida, Himoru; Nuno, Tatsuli

PATENT ASSIGNEE(S): Sumitomo Chemical Co., Ltd.

SOURCE: Ger. Offen., 53 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PA	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE	2331900	A1	19740124	DE 1973-2331900	19730622 <
DE	2331900	C2	19861113		
JP	49020129	A	19740222	JP 1972-63507	19720623 <
JP	49020130	A	19740222	JP 1972-63508	19720623 <
JP	51024491	В	19760724		
JP	49020131	A	19740222	JP 1972-63512	19720623 <
JP	51040061	В	19761101		
JP	49020128	A	19740222	JP 1972-63513	19720623 <
JP	55008501	В	19800304	JP 1972-63509	19720623 <
JP	49020133	A	19740222		
JP	49024924	A	19740305	JP 1972-64714	19720627 <
JP	55010577	В	19800317		
JP	49024923	A	19740305	JP 1972-66286	19720701 <
JP	49102629	A	19740927	JP 1973-18191	19730213 <
JP	52005486	В	19770214		
US	3920739	A	19751118	US 1973-378059	19730711 <

A 19720623 <--PRIORITY APPLN. INFO.: JP 1972-63507 JP 1972-63508 A 19720623 <--JP 1972-63509 A 19720623 <--JP 1972-63512 A 19720623 <--JP 1972-63513 A 19720623 <--JP 1972-64714 A 19720627 <--JP 1972-66286 A 19720701 <--JP 1973-18191 A 19730213 <--

GI For diagram(s), see printed CA Issue.
AB Four diamines I (R = H, Me; R1 = Et, CH2CH2OMe, CH2CH2NHSO2Me) were prepared in \$\frac{1100\text{ yield by alkylation of II and subsequent reaction with RiNHEt or

in S100% yield by alkylation of II and subsequent reaction with RINHEt or successively with EtNH2 and R1X(X = Cl, OH), or with EtNH2, C1S02Me, and HOCH2CH2NH2 followed by reduction over Pd/C.

HOCH2CH2NH2 followed by reduction over Pd/C. II 52177-13-6P 52177-25-GP

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 52177-13-6 ZCAPLUS

CN Ethanol, 2-[ethyl(3-methyl-4-nitrophenyl)amino]- (CA INDEX NAME)

RN 52177-25-0 ZCAPLUS

CN Benzenamine, N-ethyl-N-(2-methoxyethyl)-3-methyl-4-nitro- (CA INDEX NAME)

L70 ANSWER 44 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1974:61050 ZCAPLUS <u>Full-text</u>
DOCUMENT NUMBER: 80:61050

DOCUMENT NUMBER: 80:61050

ORIGINAL REFERENCE NO.: 80:9905a,9908a

TITLE: Steric effects in di- and triarylmethane dyes. XI.

Electronic absorption spectra of derivatives of Michler's hydrol blue, crystal violet, and malachite

green containing o-trifluoromethyl groups

AUTHOR(S): Grocock, D. E.; Hallas, G.; Hepworth, J. D. CORPORATE SOURCE: Dep. Sci., North Lindsay Coll. Technol., Sci

Dep. Sci., North Lindsay Coll. Technol., Scunthorpe, UK

SOURCE: Journal of the Chemical Society, Perkin Transactions

2: Physical Organic Chemistry (1972-1999) (1973),

(13), 1792~6

CODEN: JCPKBH: ISSN: 0300-9580

DOCUMENT TYPE:

Journal English

LANGUAGE:

AB The effects of substitution of o-CF3 groups in Michler's hydrol blue [14844-71-4], crystal violet [548-62-9], and malachite green [569-64-2] were examined Bathochromic shifts of the first bands were observed for electronically sym. dves. Addition of a CF3 group at the 2'-position of malachite green produced a conformational change which caused a hypsochromic shift of the first band. In certain cases, the combined steric and electronic effects of the CF3 groups were enough to shift the equilibrium between dve base and univalent dve cation in acid solution in favor of the former species.

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 51332-25-3 ZCAPLUS

CN Benzonitrile, 4-(dimethylamino)-2-(trifluoromethyl)- (CA INDEX NAME)



L70 ANSWER 45 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1973:478429 ZCAPLUS Full-text

DOCUMENT NUMBER:

79:78429

ORIGINAL REFERENCE NO.: 79:12713a,12716a

Antiandrogenic substituted anilides

PATENT ASSIGNEE(S): Scherico Ltd.

Fr. Demande, 40 pp.

SOURCE:

CODEN: FRXXBL Patent

DOCUMENT TYPE: LANGUAGE . French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
FR 2142803	A1	19730202	FR 1971-23250	19710625	<
FR 2142803	B1	19740830			
PRIORITY APPLN. INFO			FR 1971-23250	A 19710625	<

For diagram(s), see printed CA Issue.

AB Ten anilides (I; R = CHMe2, CHMeCHMe2; R1 = H, Me, Et; R2 = CF3, SOCF3, Br, C1: R3 = NO2, C1), potential antiandrogenic agents, are prepared by thirteen

standard methods. 39240-77-29

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

39240-77-2 ZCAPLUS

Butanamide, 2,3-dimethyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

L70 ANSWER 46 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1973:58091 ZCAPLUS Full-text

DOCUMENT NUMBER: 78:58091

ORIGINAL REFERENCE NO.: 78:9207a,9210a

TITLE: Substituted anilides and thioanilides
INVENTOR(S): Neri, Rudolph O.; Topliss, John G.

INVENTOR(S): Neri, Rudolph O.; Topliss, John (
PATENT ASSIGNEE(S): Scherico Ltd.

SOURCE: Ger. Offen., 73 pp.
CODEN: GWXXBX

DOCUMENT TYPE: Patent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2130450	A	19721221	DE 1971-2130450	19710619 <
DE 2130450	B2	19771013		
JP 56049884	В	19811125	JP 1971-45301	19710624 <
PRIORITY APPLN. INFO.:			DE 1971-2130450 A	19710619 <

- AB Nine anilides 3.4-RRIG6H3NR2CXR3 (I; R = Cl, F3C, F3CSO; R1 = Cl, O2N; R2 = H, Me, Et; R3 = CHMe2, CHMeCHMe2; X = O, S), useful antiandrogenic agents, were prepared mainly by acylation of 3,4-RRIG6H3NH2 (II). Thus, Me2CHCOCl was heated with II (R = F3C, R1 = O2N) in pyridine for 1.5 hr to give I (R = F3C, R1 = O2N, R2 = H, R3 = CHMe2, X = O).
- IT 39240-77-2P
 - RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 39240-77-2 ZCAPLUS
- CN Butanamide, 2,3-dimethyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

DOCUMENT NUMBER: 72:121102

ORIGINAL REFERENCE NO.: 72:21755a,21758a

Antitubercular substances. XXI. Synthesis of TITLE:

2-nitrodiphenylamines

Belton, J. G.; McInerney, Mary AUTHOR(S):

CORPORATE SOURCE: Lab. Med. Res. Counc. Ireland, Trinity Coll., Dublin,

Proceedings of the Royal Irish Academy, Section B: SOURCE: Biological, Geological and Chemical Science (1970),

69(2), 21-9

CODEN: PRIBAN; ISSN: 0035-8983

DOCUMENT TYPE: Journal LANGUAGE:

English

GT For diagram(s), see printed CA Issue. AB

The title compds. (I) were prepared by heating halonitrobenzenes with anilines or cyclohexylamine, by treating nitroanilines with halobenzenes in the presence of anhydrous K2CO3, KIO4, and reduced Cu, by treating dinit robenzenes with haloanilines, or by replacing Cl with alkoxy, thioalkyl, or amido groups on a preformed diphenylamine. The following (I) were prepared (R1, R2, R, m.p., and % yield given): CN, H, Ph, 126° (alc.), -; F3C, H, C6H11, 80-1° (light petroleum), 75; F3C, H, Ph, 84-5° (alc.), 69; Et, H, Ph, 121° (MeOH), -; HO2C, H, Ph, 220° (alc.), -; Cl, H, Ph, 61° (alc.), -; Cl, Cl, Ph, 96°, 61; H, H, 4-ClC6H4, 150°, 25; Cl, H, 3-MeOC6 H4, 92-3°, 60; Cl, H, 4-MeOC6H4, 121°, 70; Cl, H, 3-EtOC6H4, 92°, 68; Cl, H, 4-EtOC6H4, 95°, 80; F, H, Ph, 53°, -; H, H, 3-MeOC6H4, 57-8° (alc.), 44; H, H, 4-MeOC6H4, 89°, 50; H, H, 3-EtOC6H4, 47.5-8.5° (alc.), 30; H, H, 4-EtOC6H4, 84°, 56; tert-Bu, H, Ph, 73-4° (alc.), -; EtO, H, Ph, 74°, 52; BuO, H, Ph, 57-8° (alc.), 40; EtO, H, 4-ClC6H4, 68-9° (alc.), -; Cl, H, 4-MeOC6H4, 112-14° (alc.) 40; MeO, H, 4-MeC6H4, 85-6° (alc.), 50; EtO, H, 4-MeC6H4, 75° (alc.), 50; H, H, 3-MeC6H4, 70-1° (MeOH), 71; H, Cl, 4-BrC6H4, 160-1° (C6H6-alc.), -; H, Cl, 3-ClC6H4, 109° (alc.), 30; H, Cl, 4-IC6H4, 186-7°, -; H, MeO, 4-BrC6H4, 149° (alc.), -; H, MeO, 4-IC6H4, 167° (alc.), 80-90; H, MeO, 3-C1C6H4, 107° (alc.), 80-90; C1, MeO, Ph, 103-4° (alc.), 80-90; H, MeO, C6H11, 78-9° (alc.), 80-90; H, EtO, Ph, 107-8° (alc.), -; H, MeS, 4-ClC6H4 (II), 133° (alc.), -; H, MeS, Ph, 135° (alc.), -; H, phthalimido, Ph, 203-4°, -; H, NH2, Ph, 126-7° (agueous alc.), -; H, NHAc, Ph, 202-3°, -; Cl, C6H11, Ph, 124-5° (light petroleum), -. Na2S2, Na2S.9H2O, and S in 95% aqueous alc. were added to 5,4'-dichloro-2nitrodiphenylamine in 95% aqueous alc. and HCONMe2 and heated on a water bath 2 hr to give 3,3'-bis(p-chloroanilino)-4,4'-dinitrophenyl disulfide, yellow, m. 190-1° (HCONMe2-alc.), which was used to prepare II.

ΙT 28096-55-1P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 28096-55-1 ZCAPLUS

CN Benzonitrile, 4-(cyclohexylamino)-3-nitro- (CA INDEX NAME)



L70 ANSWER 48 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1970:66670 ZCAPLUS Full-text DOCUMENT NUMBER: 72:66670

ORIGINAL REFERENCE NO.: 72:12151a,12154a

Herbicidal alkylaminophenylureas

Schwartz, Herbert; Skaptason, Joseph B. INVENTOR(S):

SOURCE: U.S., 4 pp. CODEN: USXXAM DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE _____ ----_____ _____ -----US 1967~626409 US 3484484 A 19691216 19670328 <---US 1967-626409 A 19670328 <--PRIORITY APPLN. INFO .:

GI For diagram(s), see printed CA Issue.

AB The title compounds (I) are useful as preemergent and postemergent herbicides. Thus, a mixture of 98 g 4-chloro-3-(trifluoromethyl)-nitrobenzene and 97 g Pr2NH in 150 ml MeOH was refluxed overnight to give 89 g II (R1 = R2 = Pr, X = CF3, Y = NO2), b0.5 86-9°, which was hydrogenated over Pd/C to give II (R1 = R2 = Pr, X = NH2), b1 85-8°, of which 14 g was dissolved in 150 ml C6H6, and 4.5 ml pyridine then 6 q Me2NCOC1 added at room temperature to give I (R1 = R2 = Pr, X = CF3, R3 = Me), m. 124-5° (C6H6-C6H14). Similarly prepared were the tabulated I. Similarly prepared were II (R1 = R2 = Et, X = CF3, Y = NO2), b1 95-8°, and II (R1 = R2 = Pr, X = C1, Y = NO2), m, 141-3°.

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

25900-35-0 ZCAPLUS

CN Benzenamine, N,N-diethyl-4-nitro-2-(trifluoromethyl)- (CA INDEX NAME)



L70 ANSWER 49 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1969:87264 ZCAPLUS Full-text

DOCUMENT NUMBER: 70:87264

ORIGINAL REFERENCE NO.: 70:16285a,16288a TITLE:

Nitro(trifluoromethyl)toluidides INVENTOR(S): Baker, Joseph W.

PATENT ASSIGNEE(S): Monsanto Co. SOURCE: U.S., 5 pp.

CODEN: USXXAM DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

DATENT NO

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3426049 PRIORITY APPLN. INFO.:	A	19690204	US 1965-497437 US 1965-497437 A	19651018 < 19651018 <

- AB The title compds, possessing useful microbiol, activity were prepared Thus, 0.055 mole valeryl chloride was added to 0.05 mole \alpha, \alpha, \alpha-trifluoro-4-nitro-mtoluidine (I). The mixture was refluxed until the evolution of HCl ceased, 75 ml. methylcvclohexane added, and the mixture cooled to yield α, α, α -trifluoro-4'-nitro-m-valerotoluidide (IIa), m. 94-5°. Similarly prepared were RCONHC6H3(CF3)NO2-3,4 (RCO and m.p. given): isovaleryl, (IIb), 103-4°; 2methylbutyryl, 111-12°; 4-methylvaleryl, 85-7°; hexanoyl (IIc), 73-4°; neoheptanov1, 110.5-12°; decanov1, 77-8°; undecanov1, 82-3°; cyclopentanecarbonyl, 143-4°; cyclohexanecarbonyl, 135-6°; hydrocinnamoyl, 90.5-92°; sorboyl, 123-4°; 3-ethyl-2-hexanoyl, 90-1°; 2-nonenoyl, 59-60°; 2bromononanov1, 102-3°; 2-chlorononanov1, 55-6°; nonanov1 (II), 69-70°; heptanoyl (III), 64-5°; octanoyl (IV), 72-3°; lauroyl, 78-9°; 10-undecenoyl, 77-8°: 2-nonynovl, 47-8°. A solution of 0.05 mole nonanovl chloride in 25 ml. ether was added to a solution of 0.05 mole I and 0.05 mole Et3N in 200 ml. ether. The mixture was refluxed for 3 hrs., cooled, and Et3N.-HC1 removed to yield II-I complex, m. 84-5°. An excess of ethereal HCl was added to a solution of 10 g. II-I complex in 75 ml. ether and the toluidine-HCl removed to yield II. Similarly prepared were a, a, a-trifluoro-4-methyl-4'-nitro-mvalerotoluidide-I complex, m. 97-8°; III-I complex, m. 94-5°; IV-I complex, m. 90-1°. A solution of 0.0017 mole IIa and 0.0017 mole I in 10 ml. toluenemethylcyclohexane mixture was cooled to give IIa-I complex, m. 96-7°. Similarly prepared were IIb-I complex, m. 103-4°; IIc-I complex, m. 106-7°. A solution of 0.014 mole Br in 25 ml. CC14 was added to a solution of 0.014 mole α, α, α -trifluoro-4'-nitro-m-(2-noneno)toluidide in 100 ml. CCl4 at 0-5°, the mixture kept 6 hrs. at $0-5^{\circ}$, 1.5 q. KOAc in 10 ml. EtOH added, and the mixture refluxed 30 min, to give 2,3-dibromo-a,a,a,-trifluoro-4'-nitro-mnonanotoluidide, m. 107-8°.
- IT 10023-90-2P 10023-91-3P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 10023-90-2 ZCAPLUS
- CN Hexanamide, N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

- RN 10023-91-3 ZCAPLUS
- CN Pentanamide, 4-methyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

L70 ANSWER 50 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1967:37593 ZCAPLUS Full-text

DOCUMENT NUMBER: 66:37593

ORIGINAL REFERENCE NO.: 66:7131a,7134a

TITLE: Synthesis and bacteriostatic activity of some

nitrotrifluoromethylanilides

AUTHOR(S): Baker, Joseph Willard; Bachman, Gerald L.; Schumacher,

Ignatius; Roman, Daniel P.; Tharp, Alan L.

CORPORATE SOURCE: Org. Chems. Div., Monsanto Co., St. Louis, MO, USA SOURCE: Journal of Medicinal Chemistry (1967), 10(1), 93-5

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

AB Nitrotrifluoromethylanilides (I) were prepared either by the acid-catalyzed reaction of an anhydride with a substituted aniline or by the condensation of an acid chloride or with the aniline alone or in the presence of EtSN as the

HCl acceptor. Seven anilide-aniline complexes were formed when C5-9 acid chlorides free of α -substituents were treated with 4-nitro-3-trifluoromethylaniline in the presence of Et3M. The active structures of the nitrotrifluoromethylanilides which inhibited the activity of Staphylococcus aureus included those which were substituted in the meta and para positions of the N-phenyl ring with a nitro and trifluoromethyl group and in which the acid-derived moiety incorporates alkyl, haloakyl, cycloalkyl, alkenyl, haloalkenyl, alkylenyl, and phenethyl groups and contains 5-12 C atoms. Anilides substituted in the α position possessed a lower order of activity. Benzyl and phenoxymethyl derivs, and disubstituted and orthosubstituted derivs. were inactive. All of the complexes were derivs, of active anilides and exhibited the same order of activity on a weight basis as the anilides themselves.

IT 10023-90-2P 10023-91-3P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 10023-90-2 ZCAPLUS

CN Hexanamide, N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 10023-91-3 ZCAPLUS

Pentanamide, 4-methyl-N-[4-nitro-3-(trifluoromethyl)phenyl]- (CA INDEX CN NAME)

L70 ANSWER 51 OF 51 ZCAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1942:29161 ZCAPLUS Full-text

DOCUMENT NUMBER: 36:29161

ORIGINAL REFERENCE NO.: 36:4490c-f

TITLE: Alkanolamines. X. Intermediates of pentryl analogs. Chloronitroanilinoalkanols

AUTHOR(S): Kremer, Chester B.; Meltsner, M.

SOURCE: Journal of the American Chemical Society (1942), 64,

CODEN: JACSAT: ISSN: 0002-7863 DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

cf. C. A. 34, 6942.2. The condensations were carried out by refluxing for 4 h. the C12C6H3NO2 and twice the mol. quantity of the NH2 alc. in BuOH. 2-(Ranilino)ethanols (R given): 4-chloro-2-nitro, m. 107.5°; 5-Cl isomer, m. 116°; 3-Cl isomer, m. 78.5°: 6-Cl isomer, b2 155-7°: 6-chloro-4-nitro, m. 120°. 3-(Ranilino)-1-propanols: 4-chloro-2-nitro, m. 60°; 5-Cl isomer, m. 78.5°; 6chloro-4-nitro, m. 73°. 1-(R-anilino)-2-propanols: 4-chloro-2-nitro, m. 116.5°; 5-Cl isomer, m. 109°; 3-Cl isomer, m. 83.5°; 6-chloro-4-nitro, m. 144°. 1-(Ranilino)-2-methyl-2-propanols: 4-chloro-2-nitro, m. 121.5°; 5-Cl isomer, m. 127°; 3-C1 isomer, m. 98.5°; 6-chloro-4-nitro, m. 71.5°. 2-(4-Chloro-2-nitroanilino)-2-methyl-1-propanol, m. 122°. Reduction was effected with NaHSO3 in a weakly alkaline medium: 2-(R-2aminoanilino)ethanols: 4-C1, m. 122.5°; 5-C1, m. 104.5°; 3-C1, m. 74°; 6-C1, b2 135-7°. 3-(5-Chloro-2-aminoanilino)-1-propanol, m. 73.5°. 1-(4-Chloro-2aminoanilino)-2-propanol, m. 130°; 5-Cl isomer, m. 101.5°. 1-(4-Chloro-2aminoanilino)-2-methyl-2-propanol, m. 121°; 2-(4-chloro-2-aminoanilino)-2-

methyl-1-propanol. 854666-45-8P, 2-Propanol, 1-(2-chloro-4-nitroanilino)-2-methyl-ΙT RL: PREP (Preparation)

(preparation of)

RN 854666-45-8 ZCAPLUS

CN 2-Propanol, 1-[(2-chloro-4-nitrophenyl)amino]-2-methyl- (CA INDEX NAME)

L1

L2

L3

=> d his full

(FILE 'HOME' ENTERED AT 08:38:37 ON 12 MAR 2009)

FILE 'REGISTRY' ENTERED AT 08:39:03 ON 12 MAR 2009

FILE 'ZCAPLUS' ENTERED AT 08:39:33 ON 12 MAR 2009 ACT BAS012APP/A

1 SEA SPE=ON ABB=ON PLU=ON US2005-560012 /AP

FILE 'REGISTRY' ENTERED AT 08:39:39 ON 12 MAR 2009 ACT BAS012APPRNS/A

> 186 SEA SPE=ON ABB=ON PLU=ON (101130-93-2/BI OR 106-95-6/BI OR 106-96-7/BI OR 107-08-4/BI OR 107-10-8/BI OR 109-89-7/BI OR 115416-50-7/BI OR 142-84-7/BI OR 1458-98-6/BI OR 151951-35-8/BI OR 156-87-6/BI OR 1621-24-5/BI OR 194853-86-6/BI OR 208173-21-1/BI OR 2439-54-5/BI OR 24424-99-5/BI OR 2516-47-4/BI OR 26389-60-6/BI OR 2968-33-4/BI OR 31643-49-9/BI OR 393-09-9/BI OR 393-11-3/BI OR 393-36-2/BI OR 445-02-3/BI OR 4784-77-4/BI OR 49645-18-3/BI OR 513-49-5/BI OR 51332-25-3/BI OR 534-03-2/BI OR 5813-64-9/BI OR 625-43-4/BI OR 654-70-6/BI OR 67515-59-7/BI OR 7051-34-5/BI OR 75-64-9/BI OR 762-49-2/BI OR 821776-43-6/BI OR 821776-44-7/BI OR 821776-45-8/BI OR 821776-46-9/BI OR 821776-47-0/BI OR 821776-48-1/BI OR 821776-49-2/BI OR 821776-50 -5/BI OR 821776-51-6/BI OR 821776-52-7/BI OR 821776-53-8/BI OR 821776-54-9/BI OR 821776-55-0/BI OR 821776-56-1/BI OR 821776-57 -2/BI OR 821776-58-3/BI OR 821776-59-4/BI OR 821776-60-7/BI OR 821776-61-8/BI OR 821776-62-9/BI OR 821776-63-0/BI OR 821776-64 -1/BI OR 821776-65-2/BI OR 821776-66-3/BI OR 821776-67-4/BI OR 821776-68-5/BI OR 821776-69-6/BI OR 821776-70-9/BI OR 821776-71 -0/BI OR 821776-72-1/BI OR 821776-73-2/BI OR 821776-74-3/BI OR 821776-75-4/BI OR 821776-76-5/BI OR 821776-77-6/BI OR 821776-78 -7/BI OR 821776-79-8/BI OR 821776-80-1/BI OR 821776-81-2/BI OR 821776-82-3/BI OR 821776-83-4/BI OR 821776-84-5/BI OR 821776-85 -6/BI OR 821776-86-7/BI OR 821776-87-8/BI OR 821776-88-9/BI OR 821776-89-0/BI OR 821776-90-3/BI OR 821776-91-4/BI OR 821776-92 -5/BI OR 821776-93-6/BI OR 821776-94-7/BI OR 821776-95-8/BI OR 821776-96-9/BI OR 821776-97-0/BI OR 821776-98-1/BI OR 821776-99 -2/BI OR 821777-00-8/BI OR 821777-01-9/BI OR 821777-02-0/BI OR 821777-03-1/BI OR 821777-04-2/BI OR 821777-05-3/BI OR 821777-06 -4/BI OR 821777-07-5/BI OR 821777-08-6/BI OR 821777-09-7/BI OR

ACT BAS012HITRNS/A

82177

¹⁵¹ SEA SPE=ON ABB=ON PLU=ON (101130-93-2/BI OR 115416-50-7/BI OR 151951-35-8/BI OR 49645-18-3/BI OR 3332-25-3/BI OR 821776-43-6/BI OR 821776-44-7/BI OR 821776-45-8/BI OR 821776-46-9/BI OR 821776-47-0/BI OR 821776-46-1/BI OR 821776-49-2/BI OR 821776-59-5/BI OR 821776-51-6/BI OR 821776-59-5/BI OR 821776-53-6/BI OR 821776-59-6/BI OR 821776-59-6/BI OR 821776-59-6/BI OR 821776-59-7/BI OR 821776-50-7/BI OR 821776-59-7/BI OR 821776-50-7/BI OR 821776-50-7/BI OR 821776-50-7/BI OR 821776-50-7/BI OR 821776-61-8/BI OR 821776-65-9/BI OR 821776-63-0/BI OR 821776-61-8/BI OR 821776-65-9/BI OR 821776-67-9/BI OR 821776-69-9/BI OR 821776-9/BI OR 821776-69-9/BI OR 821776-79/BI OR 821776-69-9/BI OR 821776-69-9/BI OR 821776-9/BI OR 821776-69-9/BI OR 821776-9/BI OR 82

821776-71-0/BI OR 821776-72-1/BI OR 821776-73-2/BI OR 821776-74 -3/BI OR 821776-75-4/BI OR 821776-76-5/BI OR 821776-77-6/BI OR 821776-78-7/BI OR 821776-79-8/BI OR 821776-80-1/BI OR 821776-81 -2/BI OR 821776-82-3/BI OR 821776-83-4/BI OR 821776-84-5/BI OR 821776-85-6/BI OR 821776-86-7/BI OR 821776-87-8/BI OR 821776-88 -9/BI OR 821776-89-0/BI OR 821776-90-3/BI OR 821776-91-4/BI OR 821776-92-5/BI OR 821776-93-6/BI OR 821776-94-7/BI OR 821776-95 -8/BI OR 821776-96-9/BI OR 821776-97-0/BI OR 821776-98-1/BI OR 821776-99-2/BI OR 821777-00-8/BI OR 821777-01-9/BI OR 821777-02 -0/BI OR 821777-03-1/BI OR 821777-04-2/BI OR 821777-05-3/BI OR 821777-06-4/BI OR 821777-07-5/BI OR 821777-08-6/BI OR 821777-09 -7/BI OR 821777-10-0/BI OR 821777-11-1/BI OR 821777-12-2/BI OR 821777-13-3/BT OR 821777-14-4/BT OR 821777-15-5/BT OR 821777-16 -6/BI OR 821777-17-7/BI OR 821777-18-8/BI OR 821777-19-9/BI OR 821777-20-2/BI OR 821777-21-3/BI OR 821777-22-4/BI OR 821777-23 -5/BI OR 821777-24-6/BI OR 821777-25-7/BI OR 821777-26-8/BI OR 821777-27-9/BI OR 821777-28-0/BI OR 821777-29-1/BI OR 821777-30 -4/BI OR 821777-31-5/BI OR 821777-32-6/BI OR 821777-33-7/BI OR

```
821777-34-8/BI OR 821777-35-9/BI OR 821777-36-0/BI OR 82177
              ACT BAS0120LDRNS/A
L4 (
            1) SEA SPE=ON ABB=ON PLU=ON 49645-18-3
            1) SEA SPE=ON ABB=ON PLU=ON 51332-25-3
1.5 (
L6 (
            1) SEA SPE=ON ABB=ON PLU=ON 96795-43-6
L7 (
            1) SEA SPE=ON ABB=ON PLU=ON 101130-93-2
L8 (
            1) SEA SPE=ON ABB=ON PLU=ON 115416-50-7
L9 (
            1) SEA SPE=ON ABB=ON PLU=ON 151951-35-8
L10
            6 SEA SPE=ON ABB=ON PLU=ON (L4 OR L5 OR L6 OR L7 OR L8 OR L9)
               D SCA
    FILE 'STNGUIDE' ENTERED AT 08:41:38 ON 12 MAR 2009
    FILE 'REGISTRY' ENTERED AT 08:45:14 ON 12 MAR 2009
L*** DEL 5594 S MG/ELS (P) BR/ELS
L*** DEL
             1 S ?GRIGNARD?/CNS
               D SCA
     FILE 'STNGUIDE' ENTERED AT 08:51:02 ON 12 MAR 2009
     FILE 'REGISTRY' ENTERED AT 09:22:39 ON 12 MAR 2009
               D SCA L10
L11
             3 SEA SPE=ON ABB=ON PLU=ON L10 AND X/ELS
    FILE 'ZCAPLUS' ENTERED AT 09:23:07 ON 12 MAR 2009
1.12
            12 SEA SPE=ON ABB=ON PLU=ON L11
    FILE 'REGISTRY' ENTERED AT 09:23:40 ON 12 MAR 2009
L13
               STRUCTURE UPLOADED
L14
               SCREEN 616
L15
          151 SEA SPE=ON ABB=ON PLU=ON L11 OR L3
            0 SEA SPE=ON ABB=ON PLU=ON L15 AND C>21
L16
              SCREEN 1951
1.18
            3 SEA SSS SAM L13 AND L14
L19
            6 SEA SSS SAM L13 AND (L14 NOT L17)
               D SCA
```

FILE 'STNGUIDE' ENTERED AT 09:31:23 ON 12 MAR 2009

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FILE 'REGISTRY' ENTERED AT 09:46:40 ON 12 MAR 2009
               D SCA L19
    FILE 'STNGUIDE' ENTERED AT 09:52:26 ON 12 MAR 2009
    FILE 'REGISTRY' ENTERED AT 10:01:51 ON 12 MAR 2009
L20
              STRUCTURE UPLOADED
L21
             0 SEA SSS SAM L20
L22
             1 SEA SSS SAM L20 AND (L14 NOT L17)
               D SCA
1.23
               SCREEN 2043
L24
               SCREEN 2043 OR 1951
L25
             2 SEA SSS SAM L20 AND (L14 NOT L24)
              D SCA
L26
              STRUCTURE UPLOADED
1.27
               SCREEN 1993
L28
             1 SEA SSS SAM L26 AND ((L14 AND L27) NOT L24)
               D SCA
L29
             0 SEA SPE=ON ABB=ON PLU=ON L3 AND NRS>2
L30
            29 SEA SPE=ON ABB=ON PLU=ON L3 AND 2/NRS
               D SCA L30
L31
               SCREEN 1840
L32
               SCREEN 1840 OR 2043 OR 1951
L33
             2 SEA SSS SAM L26 AND ((L14 AND L27) NOT L32)
               D SCA
L34
               ANALYZE PLU=ON L30 1- RID : 4 TERMS
               D
L35
        898352 SEA SPE=ON ABB=ON PLU=ON (N>1 AND ((46.150.18/RID AND
               1/NRS) OR (46.150.18/RID AND (1.13.1/RID OR 16.127.1/RID OR
               46.150.1/RID) AND 2/NRS))) NOT PMS/CI
             7 SEA SUB=L35 SSS SAM L26 AND ((L14 AND L27) NOT L32)
L36
               D SCA
L37
             8 SEA SUB=L3 SSS SAM L26 AND ((L14 AND L27) NOT L32)
               D SCA
          1054 SEA SUB=L35 SSS FUL L26 AND ((L14 AND L27) NOT L32)
L38
               SAVE TEMP L38 BAS012STR26L/A
    FILE 'ZCAPLUS' ENTERED AT 10:25:21 ON 12 MAR 2009
L39
          3024 SEA SPE=ON ABB=ON PLU=ON L38
    FILE 'REGISTRY' ENTERED AT 10:26:04 ON 12 MAR 2009
L40
          1054 SEA SPE=ON ABB=ON PLU=ON L38 AND C<22
1.41
           143 SEA SPE=ON ABB=ON PLU=ON L3 AND C>10
L42
             8 SEA SPE=ON ABB=ON PLU=ON L3 NOT L41
               D SCA
L43
           994 SEA SPE=ON ABB=ON PLU=ON L38 AND C>8
    FILE 'ZCAPLUS' ENTERED AT 10:28:08 ON 12 MAR 2009
          2557 SEA SPE=ON ABB=ON PLU=ON L43
L44
    FILE 'REGISTRY' ENTERED AT 10:28:34 ON 12 MAR 2009
L45
            97 SEA SPE=ON ABB=ON PLU=ON L3 AND F/ELS
            54 SEA SPE=ON ABB=ON PLU=ON L3 NOT L45
L46
L47
           135 SEA SPE=ON ABB=ON PLU=ON L38 AND L3
            16 SEA SPE=ON ABB=ON PLU=ON L3 NOT L47
L48
              D SCA
L49
             O SEA SPE=ON ABB=ON PLU=ON L47 AND BR/ELS
L50
             1 SEA SPE=ON ABB=ON PLU=ON L48 AND BR/ELS AND 9/F
```

D SCA

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10/560012
L51
           10 SEA SPE=ON ABB=ON PLU=ON L48 NOT BR/ELS
               D SCA
1.52
             6 SEA SPE=ON ABB=ON PLU=ON L51 AND F/ELS
L53
             7 SEA SPE=ON ABB=ON PLU=ON L50 OR L52
L54
           142 SEA SPE=ON ABB=ON PLU=ON L47 OR L53
               D COST
               SEL RN
               SEL MF
               D COST FULL
L55
         65815 SEA SPE=ON ABB=ON PLU=ON (C14H19N3O2/MF OR C13H17N3O2/MF OR
               C12H13F3N2/MF OR C13H12F6N2O/MF OR C13H13N3O2/MF OR C15H19F3N2/
               MF OR C11H11F3N2O2/MF OR C11H8F6N2/MF OR C12H11F3N2/MF OR
               C13H15F3N2/MF OR C13H15N3O2/MF OR C13H16N2O2/MF OR C13H17CLN2/M
               F OR C13H17N3O3/MF OR C13H20N2O2/MF OR C14H14F6N2/MF OR
               C14H15F3N2O/MF OR C14H17F3N2/MF OR C14H17F3N2O2/MF OR C14H17N3O
               2/MF OR C15H17F3N2/MF OR C15H17F3N2O/MF OR C15H17N3O2/MF OR
               C15H19F3N2O/MF OR C15H19F3N2O2/MF OR C16H17F3N2/MF OR C16H21F3N
               2/MF OR C10H11F3N2O2/MF OR C10H11F3N2O3/MF OR C10H13CLN2O3/MF
               OR C10H6F6N2/MF OR C10H9F3N2/MF OR C11H11F3N2/MF OR C11H11F3N2O
               /MF OR C11H13F3N2O2/MF OR C11H13F3N2O3/MF OR C11H13N3O3/MF OR
               C11H15CLN2O2/MF OR C11H16CLN3O2/MF OR C11H16N2O3/MF OR
               C11H7BRF9N/MF OR C12H10F6N2/MF OR C12H10F6N2O/MF OR C12H11F5N2/
               MF OR C12H11F6N3/MF OR C12H13F3N2O/MF OR C12H16CLN3/MF OR
               C12H18CLN3O2/MF OR C12H18N2O3/MF OR C12H7F9N2/MF OR C12H8F8N2/M
               F OR C12H9F6N5/MF OR C12H9F7N2/MF OR C13H12F6N2/MF OR C13H12F6N
               203S/MF OR C13H12F6N2S/MF OR C13H13CLN2/MF OR C13H13F3N2/MF OR
               C13H13F3N2O2/MF OR C13H13F6N3O2S/MF OR C13H15CLN2/MF OR
               C13H15F3N2O2/MF OR C13H15F3N2O3/MF OR C13H17CLN2O/MF OR
               C13H17F3N2O2/MF OR C13H17F3N2O3/MF OR C13H17F3N2O4/MF OR
               C13H18N2O3/MF OR C13H20N2O3/MF OR C13H8F6N2/MF OR C14H11F9N2O/M
               F OR C14H12F6N2/MF OR C14H13F3N2/MF OR C14H13F6N3O/MF OR
               C14H13F6N3O2/MF OR C14H14F6N2O/MF OR C14H15F3N2/MF OR C14H17CLN
               2/MF OR C14H17F3N2O/MF OR C14H17F3N2O3/MF OR C14H17N3/MF OR
               C14H19CLN2/MF OR C14H19F3N2O2/MF OR C14H20N2O2/MF OR C14H22N2O2
               /MF OR C15H14F6N2/MF OR C15H16F6N2/MF OR C15H17F3N2O2/MF OR
               C15H19N3O2/MF OR C15H21CLN2/MF OR C15H21CLN2O2/MF OR C15H21F3N2
               02/MF OR C15H21N3O2/MF OR C15H24N2O2/MF OR C16H19F3N2/MF OR
               C16H19F3N2O2/MF OR C16H20F3NO/MF OR C16H21F3N2O/MF OR C16H21F3N
               202/MF OR C17H12F9N/MF OR C17H19F6N302/MF OR C17H23F3N2/MF OR
               C17H25N3O2/MF OR C18H27F3N4/MF OR C21H31F3N2OSI/MF OR C9H6F6N2O
               2/MF)
L56
           359 SEA SPE=ON ABB=ON PLU=ON L55 AND L38
    FILE 'ZCAPLUS' ENTERED AT 10:56:37 ON 12 MAR 2009
            68 SEA SPE=ON ABB=ON PLU=ON L56
1.58
            49 SEA SPE=ON ABB=ON PLU=ON L57 AND P/DT
L59
            19 SEA SPE=ON ABB=ON PLU=ON L57 NOT L58
            16 SEA SPE=ON ABB=ON PLU=ON L59 AND PY<2004
1.60
L61
            28 SEA SPE=ON ABB=ON PLU=ON L58 AND PD<20030610
            35 SEA SPE=ON ABB=ON PLU=ON L58 AND PRD<20030610
L62
L63
            30 SEA SPE=ON ABB=ON PLU=ON L58 AND AD<20030610
            51 SEA SPE=ON ABB=ON PLU=ON (L60 OR L61 OR L62 OR L63)
L64
               D COST
L65
             1 SEA SPE=ON ABB=ON PLU=ON L54 AND L1
```

FILE 'REGISTRY' ENTERED AT 10:59:42 ON 12 MAR 2009

142 SEA SPE=ON ABB=ON PLU=ON L55 AND L54

366 SEA SPE=ON ABB=ON PLU=ON L56 OR L66
FILE 'ZCAPLUS' ENTERED AT 11:00:41 ON 12 MAR 2009

L66

L67

```
L68 68 SEA SPE=ON ABB=ON PLU=ON L67
L69 1 SEA SPE=ON ABB=ON PLU=ON L68 AND L1
L70 51 SEA SPE=ON ABB=ON PLU=ON L68 AND L64
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FILE 'REGISTRY' ENTERED AT 11:02:00 ON 12 MAR 2009

FILE 'ZCAPLUS' ENTERED AT 11:02:04 ON 12 MAR 2009 D STAT QUE L69 D IBIB ABS HITSTR L69 1

FILE 'REGISTRY' ENTERED AT 11:02:42 ON 12 MAR 2009
D STAT OUE L70

FILE 'ZCAPLUS' ENTERED AT 11:02:50 ON 12 MAR 2009 D IBIB ABS HITSTR L70 1-51

FILE HOME

FILE REGISTRY

Property values tagged with IC are from the ${\tt ZIC/VINITI}$ data file provided by ${\tt InfoChem.}$

STRUCTURE FILE UPDATES: 11 MAR 2009 HIGHEST RN 1119363-64-2 DICTIONARY FILE UPDATES: 11 MAR 2009 HIGHEST RN 1119363-64-2

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FILE ZCAPLUS

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FILE COVERS 1907 - 12 Mar 2009 VOL 150 ISS 11 FILE LAST UPDATED: 11 Mar 2009 (20090311/ED)

ZCAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE STNGUIDE

FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: Mar 6, 2009 (20090306/UP).

=>

chain nodes : 8 9 10 11 15 20 29 30 31 32 33 35 37 38 39 41 49 50 51 52 53 54 55 56 57 58 66 67 68 69 70 71 72 76 77 78 79 87 88 89 90 91 92 93 94 95 99 100 101 102 103 104 106 107 108 109 110 111 112 113 115 116 117 118 120 121 122 123 124 125 135 136 137 138 139 140 143 144 145 146 150 153 154 156 ring nodes : 1 2 3 4 5 6 22 23 24 25 26 27 43 44 45 46 47 48 60 61 62 63 64 65 73 74 75 81 82 83 84 85 86 126 127 128 chain bonds : $1-20 \quad 2-32 \quad 3-31 \quad 4-8 \quad 5-35 \quad 6-33 \quad 9-153 \quad 9-154 \quad 15-156 \quad 22-30 \quad 23-39 \quad 24-37 \quad 25-29 \quad 9-154 \quad 15-156 \quad 20-30 \quad 20-39 \quad 2$ 26-38 27-41 43-54 44-50 45-49 46-52 47-53 48-51 54-55 54-56 55-58 56-57 60-70 61-76 62-69 63-66 64-78 65-77 66-67 66-68 70-71 70-72 72-73 79-84 81-88 82-94 83-95 85-87 86-93 88-90 88-89 89-92 90-91 100-101 101-103 102-103 102-104 106-108

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10/560012
106-107 109-110
110-111 110-113 111-112 115-150 116-117 117-118 117-120 121-122 121-123
124-125 135-136
136-137 138-139 139-140 143-144 143-145 143-146
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6 22-23 22-27 23-24 24-25 25-26 26-27 43-44 43-
48
44-45 45-46 46-47 47-48 60-61 60-65 61-62 62-63 63-64 64-65 73-74 73-75
74-75 81-82
81-86 82-83 83-84 84-85 85-86 126-127 126-128 127-128
exact/norm bonds :
1-20 4-8 5-35 9-153 9-154 15-156 22-30 25-29 27-41 43-54 60-70 66-67
73-74 73-75 74-75 81-88 106-107 110-113 115-150 116-117 117-118 117-120
126-127 126-128
127-128 135-136 136-137 138-139 139-140 143-144 143-145 143-146
exact bonds :
2-32 3-31 6-33 23-39 24-37 26-38 44-50 45-49 46-52 47-53 48-51 54-55
54-56 55-58 56-57 61-76 62-69 63-66 64-78 65-77 66-68 70-71 70-72 72-73
79-84 82-94 83-95
85-87 86-93 88-90 88-89 89-92 90-91 100-101 101-103 102-103 102-104 106-
108 109-110
110-111 111-112 121-122 121-123 124-125
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6 22-23 22-27 23-24 24-25 25-26 26-27 43-44 43-
48
44-45 45-46 46-47 47-48 60-61 60-65 61-62 62-63 63-64 64-65 81-82 81-86
82-83 83-84
84-85 85-86
isolated ring systems :
containing 1 : 22 : 43 : 60 : 81 :
G1:CN, NO2
G3:[*1],[*2]
G4:CF3,C1,NO2,CH3,OH,CN
G5:CH3,CF3,NO2,C1
G6:[*3],[*4],[*5],[*6],[*7]
G7:CH3,MeO,t-BuO
G8:CF3,OH,MeO,EtO,NH2,[*8],[*9],[*10],[*11],[*12],[*13],[*14],[*15]
G9:[*16],[*17],[*18],[*19],[*20]
Hydrogen count :
9:= exact 0 15:= exact 1
Connectivity :
9:3 E exact RC ring/chain 10:1 E exact RC ring/chain 11:1 E exact RC ring/chain
15:2 E exact RC ring/chain 79:1 E exact RC ring/chain 115:2 E exact RC ring/chain
125:2 E exact
RC ring/chain 126:2 E exact RC ring/chain 128:2 E exact RC ring/chain 138:2 E
exact RC ring/chain
```

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 8:CLASS 9:CLASS 10:CLASS 11:Atom 15:CLASS 20:CLASS 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:Atom 29:CLASS

Match level :

10/560012 30:CLASS 31:CLASS 32:CLASS 33:CLASS 35:CLASS 37:CLASS 38:CLASS 39:CLASS 41:CLASS 43:Atom 44:Atom 45:Atom 46:Atom 47:Atom 48:Atom 49:CLASS 50:CLASS 51:CLASS 52:CLASS 53:CLASS 54:CLASS 55:CLASS 56:CLASS 57:CLASS 58:CLASS 60:Atom 61:Atom 62:Atom 63:Atom 64:Atom 65:Atom 66:CLASS 67:CLASS 68:CLASS 69:CLASS 70:CLASS 71:CLASS 72:CLASS 73:Atom 74:Atom 75:Atom 76:CLASS 77:CLASS 78:CLASS 79:Atom 81:Atom 82:Atom 83:Atom 84:Atom 85:Atom 86:Atom 87:CLASS 88:CLASS 89:CLASS 90:CLASS 91:CLASS 92:CLASS 93:CLASS 94:CLASS 95:CLASS 99:CLASS 100:CLASS 101:CLASS 102:CLASS 104:CLASS 106:CLASS 107:CLASS 108:CLASS 109:CLASS 110:CLASS 111:CLASS 112:CLASS 113:CLASS 115:CLASS 116:CLASS 117:CLASS 118:CLASS 120:CLASS 121:CLASS 122:CLASS 123:CLASS 124:CLASS 125:CLASS 126:Atom 127:Atom 128:Atom 135:CLASS 136:CLASS 137:CLASS 138:CLASS 139:CLASS 140:CLASS 143:CLASS 144:CLASS 145:CLASS 146:CLASS 150:CLASS 153:CLASS 154:CLASS 156:CLASS Generic attributes : 11: : Saturated Saturation 79: Saturation : Unsaturated Number of Carbon Atoms : less than 7 Type of Ring System : Monocyclic Element Count : Node 10: Limited C, C7

Node 11: Limited

C, C6

Node 79: Limited C, C6 Node 115: Limited C, C6